# PHASE RELATIONS AND CERTAIN MAGNETIC CHARACTERISTICS OF SOME RE-Co-Fe ALLOYS

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INTERDISCIPLINARY PROGRAMME IN MATERIALS SCIENCE

INDIAN INSTITUTE OF TECHNOLOGY KANPUR

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## PHASE RELATIONS AND CERTAIN MAGNETIC CHARACTERISTICS OF SOME RE-Co-Fe ALLOYS

A Thesis Submitted
In Partial Fulfilment of the Requirements
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MASTER OF TECHNOLOGY

By M. V. SATYANARAYANA

to the

INDIAN INSTITUTE OF TECHNOLOGY KANPUR
JANUARY, 1977

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### CERTIFICATE

This is to certify that this work on 'Phase Relations and Certain Magnetic Characteristics of some RE-Co-Fe Alloys' has been carried out under my supervision and it has not been submitted elsewhere for a degree.

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#### SYNOPSIS

Rare Earth - Cobalt internetallic compounds represent a new class of naterials with excellent permanent magnet properties. But, the high raw naterial costs restrict the application of these magnets to limited fields. The use of relatively cheap raw naterials, mischnetal and iron to replace the individual rare earth elements and part of cobalt is very helpful from an economic point of view if a compromise with the magnetic properties can be made. With these points in view, a study of the MM-Co-Fe system has been taken up in the present investigation.

To establish the different phases that exist in the MM-Co-Fe system, the phase equilibria work was carried out. Alloys of different compositions were nelted, under purified argon atmosphere, in a non-consumable tungsten electrode water-cooled copper hearth arc nelting furnace. A gas purification system was fabricated and used for purifying the argon gas. The as cast alloys were sealed in evacuated quartz tubes (20x10<sup>-3</sup> nm Hg) and annealed at 900°C for 4 days. The annealed alloys were used for netallographic and x-ray diffraction

analysis. Nital etching reagent was found to be suitable for revealing the microstructures of most of the alloys. diffractometer and a Debye-Scherrer Camera were used for obtaining the x-ray diffraction patterns of the RE-Co-Fe alloys. Bēsides the  $\Lambda_2B_7$  type phase, two new phases,  $\Lambda$  phase and S phase, were found to exist in the investigated portion of the RE-Co-Fe system. The A2B7 phase was found to deviate from the hypothetical  $^{\Lambda}_2$ B7 quasi-binary line. The  $^{\Lambda}$  phase was found to extend along the same direction as the  $^{L}2^{B}7$  phase. The S phase was found to extend parallel to the quasi-binary A2B17 line. The A and S phases appeared to have structures. similar to that of the RE2Co17 phase. The lattice parameters of the A phase alloys were found to be insensitive to the change in Fe content whereas these of the S phase were found to increase with Fe content. These lattice parameter variations could not be explained on the basis of the available data.

In order to get a first hand information of favourable magnetic characteristics of the RE-Co-Fe alloys, the Curie temperatures and the easy axis of magnetization were determined for a few important alloy compositions. The Curie temperatures ( $T_c$ ) were determined by the induction method. The  $T_c$  values of different phases appear to vary smoothly with Fe content. Among the investigated alloys, the Curie temperatures of the

S phase alloys are the highest and those of the A phase alloys are the lowest. The easy axis of magnetization of the S phase was determined in the same way as the fibre axis determination. Magnetically aligned needles were prepared from powdered alloys and the x-ray pattern was obtained using a Weissenberg Camera. Etching of the alloy powder was found to produce good streaky pattern as expected from a well aligned specimen. The magnetic easy axis for the S phase was found to be along the c-axis. On the basis of the available data, the S phase appears to be the nost promising for permanent magnet development.

For determining the melting points of the RE-Co-Fe alloys, an already available system was converted to a thermal Since a heater element of suitable gauge analysis apparatus. was not available, a two-layer furnace with a room temperature resistance of about 3 ohms was fabricated using available Mo wire. The furnace assembly was calibrated using standard samples of known melting points. Attempts to carry out thermal analysis of the RE-Co-Fe alloys were not successful because the furnace failed during thermal analysis of RE-Co-Fe alloys. several important observations regarding the operation of the apparatus and the nelting points of the RECo, and RE2Co, phases have been made. The nelting points of the ternary A2B17 type phase near the RE-Co binary phase appears to be above 1400°C, well above that of the binary compound. The nelting point of the RECo<sub>5</sub> type phase appears to be below 1250°C.

#### CHAPPER I

#### INTRODUCTION

Permanent magnets play ar important and esserbled part in many devices whome an electromagnet cannot be used. Materials for those magnets have been developed from the hi # carbon chromium steels to almicos in the present century (1). But, more recently, another new class of magnets based on rare earth-transition metal compounds, with outstanding permanent magnet properties, have come into existence. sequence of events which led to the development of these materials for permanent majnor applications can be listed as follows: the discovery, in 1935 by Uwbein, Weiss and Trombe (2), of the ferromagnetism of Gd, the report on the saturation magnetization of Gd-Fe and Gd-Co alloys in 1959 by Nesbitt et al. (3), the notable work of Hubbard, Adams and Gilfrich in 1960 on the permanent magnet properties of GdCo, alloy (4), and the discovery of an extremely high magnetocrystalline anisotropy for YCo, in 1966 by Hoffor and Strnat (5). The slow progress in this field upto 1966 car be traced to the unavailability of pure rare earth metals in the early periods, their high costs, the lack of understanding of the parameters that influence the magnetic properties and not recognizing GdCo<sub>5</sub> as one of a family of compounds. The discovery of Hoffer and Struct, however, was followed by extensive studies to determine the parmament magnet properties of the family of rare carth-transition metal compounds.

These compounds, particularly the light rare earth (at nos. 57-63) - cobalt compounds are reported to have high remainent magnetization and coercivity. The uneful magnetic properties of SmCo<sub>5</sub> and (Sm, Pr) Co<sub>5</sub> alloys are compared with those of the conventional permanent magnetic materials in Table I.1.

Nost of todays rare earth-transition movel permanent nagmets are based on SaCo<sub>5</sub>. This is because of the fact that SaCo<sub>5</sub> magnets with excellent angmetic properties can be fabricated relatively easily. Nevertheless it has become increasingly apparent that not all the properties of a permanent magnet are equally important in a given application. For example, in applications with a relatively narrow aim gap where the magnet operates at a high load line - B/\(\mu\_0\mu\) ( like in motors and generators), on extremely high resistance to demagnetization is not necessary. As a result, in such applications, the potential of the SaCo<sub>5</sub> magnets is not completely utilized. Also, the relatively high price of same restricts the application of SaCo<sub>5</sub> magnets to fields where the costs are determined by production processes rather than

TABLE I.1

Magnetic Properties of Some Selected Permanent Magnetic Materials(1)

Material	Typica	o Magnetic Pr	onerties
Rei	enent Induct Br, KG	1 Force	Product (BH) MG0e
3.5 percent Cr steel	9•5	66	0.29
40 porcont Co steel	10.0	242	1.03
Alnico l	6.6	540	1.40
Alnico 5	12.0	720	5.0
Almico 5 DO	13.3	685	6.5
Alnico 8	7.1	2000	<b>5•5</b>
Alnico 9	10.4	1600	8.5
Cunife 1	5.7	590	1.85
Cunico 1	<b>3.</b> 4	710	0.85
Remalloy	10.5	250	1.1
Vicalloy 2	10.0	450	3.0
Platinum-Cobalt	6.45	4300	9•5
Coramic			
1) isotropic berium ferrit	c 2 <b>.25</b>	1850	1.15
ii) anisotropic barium ferr	ite3.95	2400	3 <b>.5</b>
ii) anisotropic strontium ferrite	3.425	3300	2.9
SmCo5 <sup>x</sup>	9.0	9000	20.0
(Sm,Pr)Co5X	9.9	6800	23.0

values for these materials are quoted from references(6) and (7) respectively.

by those of the raw natericl. The wider scope of the application of mare earth-cobalt magnets requires the material costs to be lowered while new types of angrets with specific application - oriented properties are developed. Magnets of somewhat inferior magnetic qualities than the SmCo, magnets may thus be useful for various applications. The rare carth olements occur together in the ores and at present it is the cost of scratation that makes the individual metals so expensive. Since the highest magnetic qualities are not always required, efforts are underway to replace samarium by the cheaper mixture of rare carths, the mischmetal (MI). of particular importance to India where vast deposits of rare earths (in Kerala sands) are available. The other component of the rare earth magnets, cobalt, is also expensive, particularly in India because it has to be imported. indications that partial iron substitution for cobalt improves the properties of cobalt rich rere earth-cobalt alloys to some extent (3). Hence, substitution of Co by iron in LEC-Co magnets, at least in part, is expected to be a valuable convribution towards cost reduction. So, a thorough investigation of the MI-Co-Fo system was thought very escential and has been taken up in this investigation.

#### I.1 Literature Revieu:

The term 'Rare Earth' refers to the 4f-transition elements with atomic numbers between 57 (Lanthanum) and 71 (Latetiam). Yttrium (39) is also included in this group because of its close resemblance in properties to the many carths. Gadolinium (64) for which the 4f shell is helf filled divided the wave carths into light (less than helf filled) and heavy (more than helf filled) rare earths. For the light rare earths the orbital moment L and the spin moment S are antiparallel to each other so that the resultant magnetic non-ent has the ragnitude J: |L-S|. On the other hand, for the heavy rare earths the orbital and spin moments are parallel and J: |L+S|. This behaviour has important implications for the total magnetic moment of the RE-Co compounds.

In their elemental form, the rare earths have very interesting regnetic properties. Some of their physical and magnetic properties are listed in Table I.2. As seen from the Table, some of the elements like Y, La or Bu have no permanent magnetic moment (in their usual valence state, +3) while others are magnetic. Due to strong crystal field effects, the ferromagnetic rare earths also exhibit extremely high magnetocrystalline anisotropy (9). The element dyspresium, for example, has the highest known saturation magnetization, 37.950 gauss, and the highest uninxial anisotropy constant,

Elements (9) Physical and Mainctic Properties of Rere Berth TABLE 142 C . I. 2

B] cmcnt	Atomic no.	Atomic Metallic no. valence	Meltir point;	Boilling point, oc	Free etom elec- tronic configura- tion.	llegnetic menent/atom VB	Curic Temp-
T.anthanun	57		920	3454	5 <u>d</u> 1682	0.	Monc
Certur	28	3,4	798	3257	S S	2.56	i
Praseodymium	59	20	SSI	327.2	ત્રે-6લ	3.62	None
Neodymj.v.n	09	20	1010	31.27	42,50,68	2•68	ì
Promothium	19	20	1035	2700		2,83	ı
Semerium	62	23	1072	1778	3	1.6	i
Buropium	63	C)	822	1597	CO.	3.45	-165
Gadolinium	64	10	1311	3233	S	76.7	<b>9T</b> +
Terbium	65	M	1360	3041	418591652	9.72	<b></b> 26
Dysprosiu"	99	20	1409	2335	4295d1682	10.6	-158
Holimium	L9	~	1/10	2720	4r105d16s2	10.6	1253
Erbium	89	20	1522	2510	d-65	9.6	1254
Thulium	ଓଡ	2	1545	1.727	19 <sub>1</sub> 9	7.6	i
Ytterbium	70	2	824	237	,4 <mark>1</mark> 65	4.5	Nonc
Lutotium	7.1	8	1.656	3315	68	0	Tone
Yttrium	39	~	1523	3337	44 <sup>4</sup> 55	0	Nonc
				A THE PARTY OF THE	er gement i styre i de segle i i de la segle segle segle segle i de segle segle i de la segle segle segle segle	general en i de la company	geraprodite og vidigalag gördi Diliprogittas pirigijalapi

L = 10.9x10<sup>8</sup> erg/cm<sup>3</sup> at 0°K. However, the nagnotic order in the RE elements occurs only at low temperatures. The highest Curio temperature, among the rare earth elements, is that of Gd, 16°C (Table I.2). So, the nure rare earths are of no use for conventional regretic applications. When a rare earth element and a 3d-transition notal are alloyed together, a number of intermediate phases form. These phases show fewro-, ferri-, or antiferroma metic order at room temperature. Some of the ferromagnetic phases show Curie temperatures of several hundred degrees centigrade. In many of these phases the rare earth elements induce a high uniaxial crystal anisotropy which makes those substances interesting as hard magnetic materials (4).

been studied for Mn, Fe, Co and Mi and magnetic properties of various phases formed have been determined to see which of the RE-transition metal phases are of practical importance. In the case of RE-Mn systems, the three phases, REMn<sub>12</sub>, RE<sub>6</sub>Mn<sub>23</sub> and REMn<sub>2</sub> (No REMn<sub>5</sub> compound exists) have been found to exhibit either antiferromagnetic order or they are ferremagnetic with Curic temperatures too low to be of interest for permanent magnet use (10). In the case of RE-Mi systems, a variety of phases (RE<sub>2</sub>Ni<sub>7</sub>, RENi<sub>5</sub>, RE<sub>2</sub>Ni<sub>17</sub> and other high RE phases) form but they have low Curic points (11).

The Ce-Fe diagram is shown in Fig. I.l. The other RE-Fe diagrams are reported to be havin; similar features (12). In the RE-Fe systems of Ce.Nd, Pr and Sm, recent investigations indicate that the RAFe, phases, which were proposed by earlier investigators, do not exist. The only two phases that exist are REFe, and RE2Fe, 7. The RE2Fe, 7 compounds are reported to have high saturation magnetization and uniaxial crystal symmetry, but they have low Curic points (T  $_{\rm c}$  < 180 ) (13). To increases but the saturation magnetization drops. RE-transition metal compounds of Mn, Fe and Mi are not useful for permanent magnet applications. The RE-Co alloys, however, show regretic characteristics (indicated later) which are highly attractive for permanent magnet applications. Also, there are indications that addition of Fe to some of the RE-Co compounds improves their magnetic characteristics (8). This, even though the RE-Fe alloys are not important for permanent magnet applications, RE-Co-Fe alloys may be quite promising.

The RE-Co systems are the most widely investigated ones among the RE-transition metal systems. Several of the binary RE-Co phase diagrams (for RE = Co, La, Pr, Nd and Sm) are given in Figs. I.1 to I.3. As shown in the figures, all the binary RE-Co systems show the existence of RE<sub>2</sub>Co<sub>7</sub>, RECo<sub>5</sub> and RE<sub>2</sub>Co<sub>17</sub> phases and several high RE-Co phases. Those high

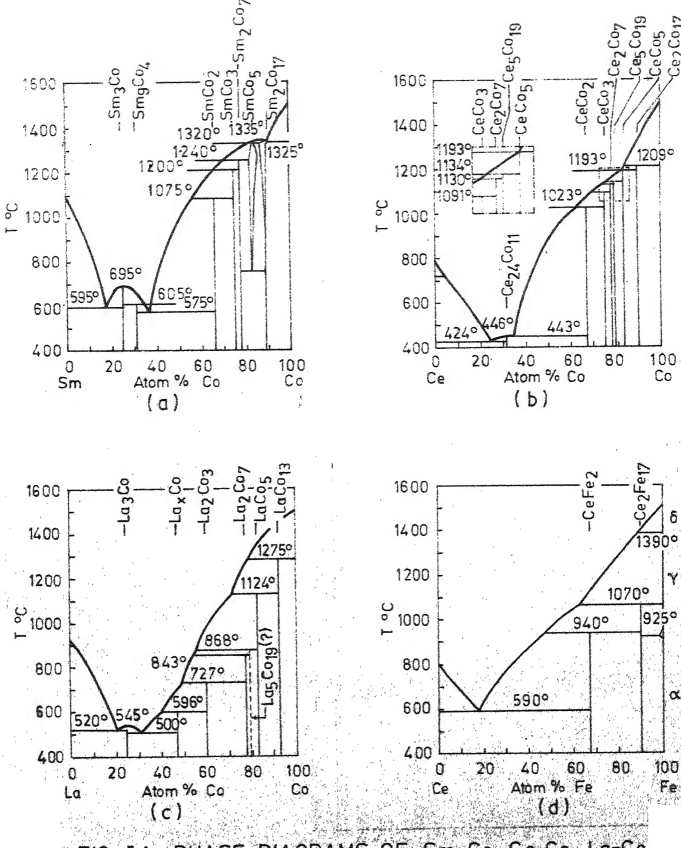


FIG. I.1. PHASE DIAGRAMS OF Sm-Co.Ce-Co.La-Co AND Ce Fe SYSTEMS(14).

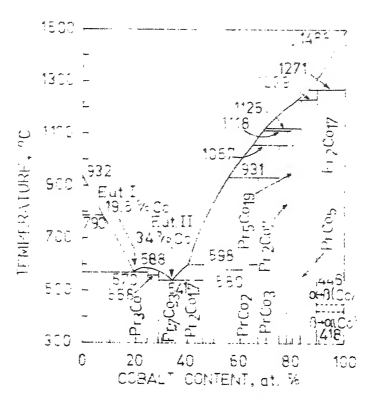


FIG. I. 2. PHASE DIAGRAM OF THE Pr-Co SYSTEM (15)

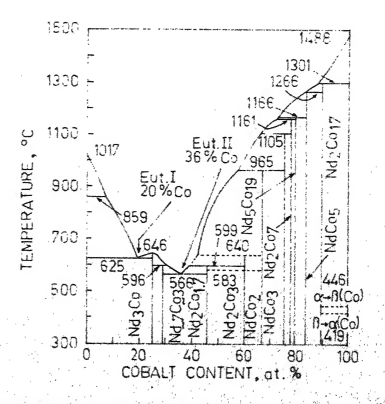


FIG I.3. PHASE DIAGRAM OF THE Nd-Co SYSTEM (15)

Co phases usually have a very narrow phase region. Only RECo, and RE2Co, 7 compounds for Ce and Sm ore reported to have an appreciable homogeneity region; the RECo $_{\kappa}$  homogeneity region extends towards higher Co concentrations whereas the RE200,7 Lomogeneity relion extends towards concentrations poorer in Co (16,17). Some of the RECo, compounds are reported to decompose extectoidally into RE2Co, and RE2Co,7 st low temperatures. For example, SaCo5 decomposes at 750°C (18). Even though Buschow (18) has reported that the other RECog phases also decompose at lew temperatures, Ray (19) could not find any evidence of it. The HE Co, type compounds usually crystalline in the hexagonal Collin type structure with the exception that La\_Co, has both bexagoned and rhombohodred polymorphic structures (20). All the RECog compounds have hexagonel CaCu, type structures whereas the RE2Co,7 compounds usually have rhombohedral Th22n17 type structures. and Co2Co17, however, have both hexagonal (high temperature) and rhombohedral The Zn, 7 type structures (12). Lattice constants and structural characteristics of some of the high Co contairing phases, RE2Co7, RECc5 and RE2Co17 are listed in Table I.3.

TABLE I.3

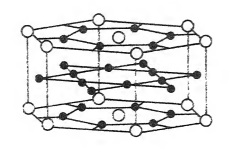
Lattice Constants and Structural Characteristics of Som Important RE-Co Phases (15,16,20,56)

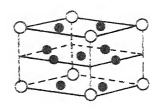
			The second secon					A THE RESIDENCE OF THE PROPERTY OF THE PROPERT
dompound		2	-		RECO <sub>5</sub>			077
El caont	<b>™</b>	4 , o	Ao Stracture type	ය . අ	A. 10	otructur.c type	d d	type our c
Gorrium .	6.6.5	24.47	Hexp_corel Co2=17 type	4.928	4.015	Hexagonal Gacu <sub>s</sub> type	8.302 8.301	8.13 Hexagenel TheN17type 12.207 Rhembehedrel The En 17type
Lanthanum	5.101	24.511	Eoxagonel Gezkiq tere	5.1	3,958	Hexagonel 11.344 Cacu <sub>s</sub> typo	11.344	- Cubic
•	5.11	96.69	Rhorbehodrel Gd <sub>2</sub> Co <sub>7</sub> t <sub>2</sub> pe					
II eo dymium	5.063	24.45	Hexagonal Gozái, tzpo	5.028	3.977	Hoxagonal CaGu <sub>5</sub> tgne	8,426	12.245 Rhombohodral The Zn Type
Prascodynium	5.072	24.57	Hexaconcl de <sub>z</sub> Hig	5.032	3.992	Ecxagenal CaGu <sub>5</sub> typo	0.436	12.276 Rholboucdrel Th $_2^{\mathrm{Zn}}_{17}$ typo
Semurium	5.041	24.327	24.327 Hexagoral Coz <sup>M</sup> 1,7°2P	5,002	3.96¢	Hexa jenel CeCu <sub>5</sub> type	8.395	1.2.216 Rhonbohodrel The Zn <sub>17</sub> typo (upto chout 120000)
							8,36	8,515 Hexagond

K

The hexagonal Cadus structure of RE-Co calloys consists of an infinite stacking sequence of two different layers of atoms in the direction of two c-axis - a hexagonal arrangement of Co atoms followed by a mixed layer of RE and Co atoms as shown in Fig. I.4(a). The hexagonal call of AEOo5 phase contains 3 rhembohodral unit calls (Fig. I.4(b)). The structures of the compounds in the range RECo2 to RE2Co17 can be derived from the hexagonal RECo5 structure (11,22). The mode of formation of hexagonal and rhembohodral structures of the RE2Co7 and RE2Co17 compounds from the RECo5 structure is schematically shown in Figs. I.5(a) and (b).

Before discussing the magnetic properties of the rare earth alloys, it will be of interest to recall which combination of properties qualifies a ferromagnetic substance for use as a permanent magnetic material. First of all, it must have a high saturation regnetization,  $K_{\rm S}$ , at room temperature so that it can produce a romanent flux  $(B_{\rm R})$  of useful magnitude. The theoretical upper limit of the energy product (the maximum value of BXH in the second quadrant of the hysterisis loop),  $(BH)_{\rm max} = (2\pi K_{\rm S})^2$  in C.G.S. units, is also determined by  $M_{\rm S}$ . Second, the Curie temperature,  $T_{\rm C}$ , must be high. Finally, it is necessary that the intrinsic coercive force be sufficiently high, namely,  $M_{\rm C} > 2\pi M_{\rm S}$ , to allow a close approach to the theoretical energy product.





O RARE EARTH . Cobalt

(a) (b)

FIG.I.4. CaCus TYPE STRUCTURE: (a) HEXAGONAL UNIT CELL, (b) RHOMBOHEDRAL UNIT CELL (14)

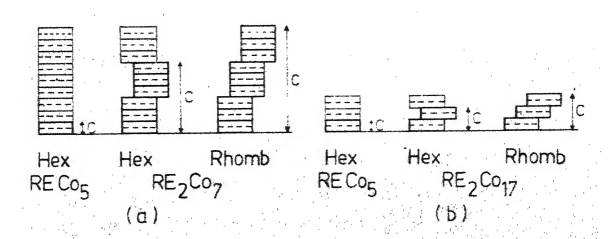


FIG. I.5. MODE OF FORMATION OF HEXAGONAL AND RHOMBOHEDRAL REXCOY ALLOYS OUT OF THE RECOS STRUCTURE: (a) RE2CO7 ALLOY, (b) RE2CO17 ALLOY.(11)

According to the modern concept of permanent magnetism, such high coercive forces are only expected of a substance which has very large magnetocrystalline anisotropy (determined by the anisotropy field,  $H_A$ ) and a single easy axis of magnetization.

The studies of Nassau, Cherry and Wallace (23) and Nesbitt et al (24) on the temperature dependence of magnetization for RECos compounds reverled two different types of behaviour. For Y and the light rare cartis, the wa netization increased with decreasing temperature whereas for Gd and the heavy rare earths, the magnetization decreased with decreasing temperature. These observations are consistent with the idea that the spin of the rare earth atoms always conclos entiperallel with that of the cobelt atoms. Thus, in the light rare earth alloys, the total moment of the RE (J = | L-S| ) adds to that of the cobalt leading to higher magnetization values. But, ir the Gd and heavy rare earth alloys, the total rare earth moment (J = |L+S|) is antiparallel to that of cobalt leading to a two sub-lattice ferrimagnetism with the associated lowering of the total magnetization. Thus, the strongest magnetic moments are observed in alloys prepared with the largest rare earths or even with the non-magnetic Y. This is shown in Fig. I.6 in which the approximate values of the saturation induction,

AnN<sub>E</sub>, for And Regron and Regron alloys are plotted as a function of the rare earth atoms. If the RB-Co magnets are to cornete with the existing almice magnets, for example, the saturation induction at room temperature of the RB-Co compounds must reach at least 6000 Gauss. From Fig. 1.6 it follows, then, that the convenient group of compounds in limited to the RE<sub>2</sub>Co<sub>17</sub> and RECo<sub>5</sub> compounds of Y and the light rare earths. Incidentally, these compounds also have high Curio temperatures as is shown in Fig. 1.7.

The magnetic properties of the useful RECo, compounds are listed in Table I.4. As indicated in the table, the RECo, phases have very large (largest known) magnetocrystalline emisotropy values (as shown by the large anisotropy fields) and substantial values of saturation mometization at room temperature. They also have the desirable casy c-axis magnetic symmetry. Permanent magnets based on these materials have now been made with proviously unatteinable ccercivities and energy products. But, only in the case of SmCog compound, energy product close to the theoretical limits has been achieved. As for example, Das (6)th has reported a (BH) wax value of 20 MGOo for liquid phase sintemed which is very close to the theoretical value. SmCo<sub>5</sub> alleys In the case of the other RECo5 compounds, further work has to be done to realize the theoretical values of (BH) max.

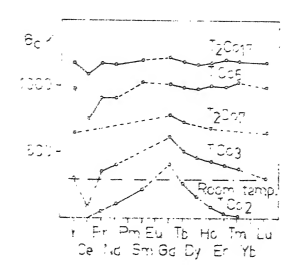


FIG. I.7. CURIE TEMPERATURES OF SEVERAL RARE EARTH(T) COBALT(Co) PHASES(25).

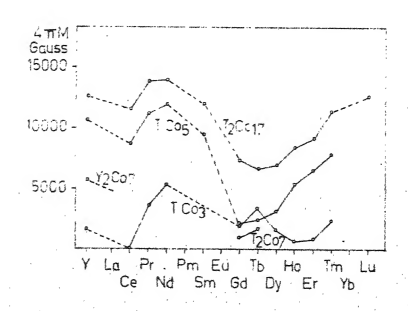


FIG. I. 6. SATURATION INDUCTION OF SEVERAL RARE EARTH(T) COBALT(Co) PHASES(25)

TABLE 1.4

Magnetic Properties of RECo<sub>5</sub> Compounds (1)

<b>೦</b> 0ಇಭಿರಾಗದೆ.	Saturation Magnetization And KG	Anisotropy Field, H <sub>A</sub> KOe	Theoretical Energy Product, (BH)
YCo <sub>5</sub>	10,6	130	28.1
Lanthenum	9.09	175	20.6
Cerius	7.7	170-210	14.8
Praseodymium	12.C	145-210	36.0
Samarium	9.65	210-290	23.0

Comperatively less amount of work has been done on the Sabrication of magnets from the RB2Co17 type compounds. Even though RE20017 compounds have large saturation nametimations and high Curis temperatures compared to the corresponding RECo, compounds (Figs. I.6 and I.7), efforts at producing waynots with good properties from the RE20017 compounds were unsuccessful. This is because of the fact that RE2Co17 compounds of light rare cartle one Y, excepting Sm2Co17, exhibit easy magnetization directions lying on the basel plane and show work or practically no anisotropy (26). S-2Co,7 exhibits cary e-exis behaviour and also has high arisotropy ( $H_A = 105 \text{ KOe}$ ) (27). Consequently, much of the world done on the RE2Co, 7 compounds has been with the Sm2Co, 7 compound. Initial results incicate that the procedure employed in indricating the RuCo, type unnets may not be applicable for the inhrication of magnets from the RE2Co, 7 compounds (8), This is probably boccuse of the fact that the  $\mathtt{RE}_2\mathtt{Co}_{17}$  type compounds are not as brittle as the RECo5 compounds and the process of powdering causes mechanical damme of the surface layers of the particles (28). At probent, investigations are going on in several laboratories abroad to find the fabrication procedure for those AE2Co,7 compounds to achieve better negretic proportics.

RECo<sub>5</sub> compounds has been studied for attaining better augmetic properties. Through these studies it is revealed that one RE can reduce another in the RECo<sub>5</sub> structure and a feel light RE additions like Md, Pr etc. improve the nagmetic characteristics of the RECo<sub>5</sub> alloys. For example, an among product of 23 MGOe has been reported for the (Sm, Pr) Co<sub>5</sub> alloy (7) whereas for the SmCo<sub>5</sub> alloy it is only 20 MGOe (6). We such studies are reported for the RE<sub>2</sub>Co<sub>17</sub> compounds.

tial real comment of cobalt by one (non-ferromagnetic) or the (one non-ferromagnetic and other ferromagnetic) transition metals of the iron group have also been investigated for realizing better me notic preparties. Extraordinarily high coercive force values have been achieved for the RE (Co<sub>1-x</sub>Cu<sub>x</sub>)<sub>5</sub> alloys by a solid state precipitation of the megnetic RECo<sub>5</sub> phase in a non-magnetic RECu<sub>5</sub> matrix. For example, as east SmCo<sub>5</sub> camples have coercivities loss than 1000 0e whereas the as east Sm(Co<sub>1-x</sub>Cu<sub>x</sub>)<sub>5</sub> alloys with x=0.3 to 0.4 have coercivities as high as 10,500 0e and it is further improved to 28,000 0e after annealing at 400°C (29). Similar improvement in coercivity has been reported for the Al substituted RECo<sub>5</sub> alloys (29). While the exercivity increases due to the addition of non-magnetic elements in NECo<sub>5</sub> phase, the

remanence is reported to decrease dristically (e.g. 9500 Gauss for SmCoCu, ) because of the dilution of the matrix by the non-magnetic copper or alluminium (29). Small additions of iron to the Cu medified SmCo<sub>5</sub> compounds, however, are reported to increase the saturation magnetization as well as coercive force. For example, the annealed (425°C, 4 hrs) compound Sm (Cu<sub>1.35</sub>Fc<sub>0.5</sub>Co<sub>3.5</sub>) showed a coercive force of 10,000 Oc which is 4000 Oc hi her than that of the corresponding compound without iron addition. The saturation induction (5900 Gauss) of the Fe containing alloy is also higher (by about 500 Gauss) than the alloy with no Fe (29).

relatively cheep and abundant 3d transition metal, iron.

-atic
But no system/investigation of the phase equilibria in the
RE-Co-Fe system has been published. Only a few workers have
investigated the limits of extension of the RECo<sub>5</sub> phase or
addition of Fe (30). From their results it appears that the
so called pseudo-binary compounds - RECo<sub>5</sub> with Fe, are
unstable. This may be because of the fact that there is no
compound corresponding to RECo<sub>5</sub> in the RE-Fe system (Fig. I.1).
Buschew has investigated a limited composition range of the
Sm-Co-Fe system (30). He observed an increase in saturation
magnetization (92 emu/g for SmCo<sub>5</sub> to 117 cmu/g for SmFeCo<sub>4.7</sub>)
and a decrease in intrinsic coercive force from 18 KOe to
10 KOe for a 1 hr. ball milled retorial. However, on

continued ball milling of the material, the coercive force was reported to increase to about 14 KOe, a value very close to that of SmCo<sub>5</sub>. Thus, the iron substituted SmCo<sub>5</sub> compounds have somewhat better magnetic properties than the compound without Fe.

No phase equilibria data is available for the substitution of Co by Fe ir RE2Co17 compounds. Ray et al (31) have mentioned that extensive solid solutions exist between RE20017 and RE2Fe17 phases at high temperatures. The --RE2(Col-xFex):7 phase, however, goes through an eutectoid transformation and at room temperature, stable RE2(Co1-Fex)17 phases are restricted only to the neighbourhood of the binary compounds. They observed that the  $RE_2(Co_{1-x}Fe_x)_{17}$  however, can be retained in a metastable state by quenching (3). Magnetic measurements were carried out with the metastable samples of the  $Sm_2(Co_{1-x}Fe_x)_{1.7}$  alloys in which x varies from 0 to 1 (32). The values of  $T_{_{\mbox{\scriptsize C}}}$ ,  $4\pi M_{_{\mbox{\scriptsize S}}}$ and  $H_A$  for these alloys are plotted as a function of x in Fig. I.8. As shown in the figure, the Curie temperatures of these alloys remain above the  $\mathbf{T_c}$  of  $\mathbf{SnCo_5}$  when Fe is substitute for Co upte about x=0.6. The saturation magnetization increases on the addition of Fe and goes through a maximum on the iron-rich side before it declines. A similar behaviour is reported for Y2(Col-xFex)17 alloys (8). This is quite analogou to the behaviour of binary Fe-Co alloys. The desirable easy c-axis behaviour is maintained upto about 50 percent iron

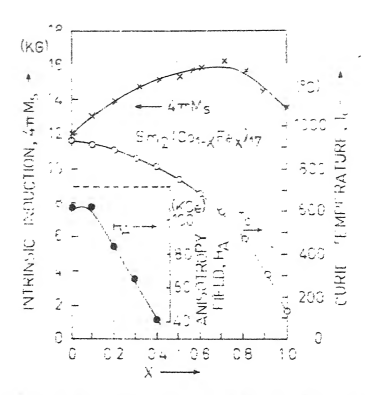


FIG I.8. VALUES OF  $4\pi M_S$ ,  $T_C$ , AND  $H_A$  FOR THE  $Sm_2(Co, Fe)_{17}$  ALLOYS (32).

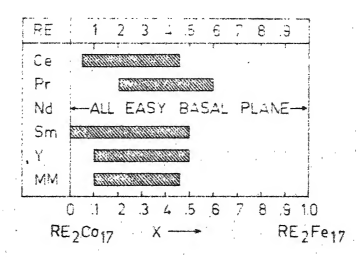


FIG 19 MAGNETIC SYMMETRY OF THE RE2(Co1-x Fex)17 PHASES(32)

substitutionin the Sm2(Co, Fe,),7 alloys, as shown in Fig. I.9. Also, the substitution of iron first enhances the easy axis anisotropy (HA) of Sm2Co,7, HA then drops, but remails attractively high upto about x=0.4. In the case of quasibinary RE2Co17 - RE2Fe17 systems of other rare carths Co. Ro and Y, the most remarkable feature of re addition is thet between 5 to 20 percent Fe substitution for Co, the c-axis bocomes the easy exis of magnetization and this is maintained upto about 45 to 60 parcers Fe substitution (Fig. I.9). The maximum values of the emisetrop, field in these cases are. however, substantially lower than those of Sm2(Co\_1-xFe\_x)17 alloys (less then 20 KOe) (32). Thus, the  $Sn_2(Co_{1-x}Fe_x)_{17}$ alloys with x=0 to 0.4 are clearly superior to the others. · Early attempts to produce A2B17 type negmets have therefore correctly concentrated on the Sm2(Co1-xFex)17 phase between 0 < x < 0.4. It is predicted that remots with (BH) max values as high as about 60 MGOs will be possible to make with the Su<sub>2</sub>(Co<sub>1-x</sub>Fc<sub>x</sub>)<sub>17</sub> alloys.

Extensive data is available on the solubility of rare cartha among themselves. Complete solid solubility in the Co-La, La-Hd and Pr-Hd systems and extended solubility for the other rare earths among themselves has been reported (33). Thus, it is expected that in FM the RE elements will be in solid solution form and when alleyed with other elements the

observed in the MM-Mg system which exactly resembles the (Ge. La, Nd) - Mg systems (34). No such data is, however, available for the MM-Co and MM-Fe systems but, it may safely be assumed that MM-transitional metal systems will show a behaviour similar to the HM-Mg system.

for the MI-Co system, several workers have investigated the magnetic properties of MMCo<sub>5</sub> phase (IM in these cases had rare carth contents of 45-60 at percent Co, 23-35 at percent La, 9-20 at percent Nd and 3-7 at percent Pr). Struct et al (35) and McCaig (36) obtained a (BH)<sub>max</sub> value of 2.3 MGOe for the field pressed MMCo<sub>5</sub> powder without sintering. Later, by using sintering techniques, Johnson and Fellows (37) and Das (38) achieved a (BH)<sub>max</sub> value of 9 MGOe for the MICo<sub>5</sub> magnets. No magnetic data is available for the corresponding MI<sub>2</sub>Co<sub>17</sub> compounds excepting that the MM<sub>2</sub>Co<sub>17</sub> compound is reported to have the easy axis of magnetization in the basal plane (26).

Small additions of other rare earths are reported to vastly improve the magnetic properties of  $(IM_{l-x}RE_{x})Co_{5}$  compounds. Several workers have investigated the system  $(IM_{l-x}Sm_{x})Co_{5}$  for x=0 to 1 and found that the magnetic properties steadily improved with Sm additions (38,39,40). For example,  $(BH)_{me,x}$  value for MMCo<sub>5</sub> is reported to be

14.5 MGOe whereas for  $MM_{0.5}$   $Sm_{0.5}$   $Co_5$ , it is reported to be 18 MGOe (39). The effects of other rare earth additions like Ce, Pr, Nd, La have also been studied. For this purpose, Hagel and Meuth (39) selected a composition (MM = RE x)0.85  $\mathrm{Sm}_{0.15}$   $\mathrm{Go}_5$  with 0 < x < 0.5. Their results can be surmarized in the following way: (1) Co reduces the  ${
m H}_{
m c}$ ,  ${
m H}_{
m A}$  and  ${
m B}_{
m R}$ .  $M_s$  is also slightly lowered; (2) La increases the  $M_c$ , slightly decreases the  ${\rm H_A}$  but does not change the  ${\rm B_R}$  and  ${\rm M_S}$ ; (3) Pr reduces the  $_{\mathbb{R}}^{\mathbb{H}}_{\mathbf{c}}$  and  $\mathbb{H}_{\mathbf{A}}$  but increases the  $\mathbf{B}_{\mathbb{R}}$  and  $\mathbf{M}_{\mathbf{s}}$ ; (4) in the case of Nd,  $_{
m M}^{
m H}_{
m c}$  is reported to increase initially (from 8110e at 0 percent Nd to 16 KOe at 10 percent Nd) but further additions of Md decrease the  $\mathbf{M}_{\mathbf{c}}$ .  $\mathbf{H}_{\mathbf{A}}$  is reported to decrease steadily with increase in Nd centent. Both  $B_{\mathrm{R}}$  and  $M_{\mathrm{s}}$  are reported to increase with No additions. From those results, Nagel and Meuth report that energy products upto 18 MGOs can be obtained with MM based magnets.

Cast permanent magnets based on MM (Co, Cu)<sub>5</sub> alloys, in which magnetic hardening is produced by solid state precipitation of a magnetic phase in a non-magnetic matrix, have also been produced and studied for their magnetic properties. Strnat et al (41) and Takata (42) obtained an energy product of 6 MGOe for the cast MM (Co,Cu)<sub>5</sub> magnets. Later, by using powder metallurgy techniques, higher energy product (\*\* 8 MGOe) magnets have been developed from

HM(Co,Cu)<sub>5</sub> alloys (43). Iron additions to NM(Co,Cu)<sub>5</sub> alloys are found to further improve the magnetic properties (44).

No magnetic data is available for both MM(Co,Fo)<sub>5</sub> and MM<sub>2</sub>(Co,Fe)<sub>17</sub> alloys. However, iron additions of about 10 percent are found to change the easy axis of ragnetization from the basal plane to the c-axis (Fig. I.9) in the MM<sub>2</sub>(Co<sub>1-x</sub>Fe<sub>x</sub>)<sub>17</sub> alloys and this c-axis magnetic symmetry is maintained upto about x=0.45.

### I.2 Statement of Problem:

The use of iron and mischmetal substantially reduces the east of RE-Co magnets even though their use may lead to somewhat inferior magnetic properties compared to those of the SmCo<sub>5</sub> magnets. However, as mentioned in section I., for certain applications, magnetic properties comparable to those of SmCo<sub>5</sub> magnetse are not really necessary. Consequently, in such applications, the NM-Co and the NM-(Co, Fe) alloys have definite economic advantage over other materials.

The ferromagnetism in transition motals Fe, Co and Mi is due to unequal filling up of the 3d sub-bands. Addition of Fe to Co (Fe having one less electron/atom than Co) causes more imbalance between the two 3d half bands of Co and as a result magnetic moment/atom increases as Fe content is increased. The moment/atom goes through a maximum near about 70 percent Fe. A similar effect has been observed with the RE-Co

systems - addition of RE appears to reduce the imbalance in the two half bands thereby lowering magnetic moment/atom of the transition metal (25). On Fe addition to RECo, and RE Co, 7 phases of Gd-(Co,Fo), Y-(Co,Fo) and Sm-(Co,Fe) systems, it has been reported (8,32,45) that magnetisation increases with increase in Fe and it passes through a maximum at a reasonably high Fe content, similar to what has been found in the Fe-Co system. On the basis of these observations, it is expected that in IM-Co-Fe system also, there will be an increase in magnetization with increase in Fe. Moreover, there may be new plases existing in the RE-Co-Fe system with magnetic properties similar or superior to the MMCo, and MM2Co, phases. With these points in view, it was thought necessary to carry out the phase equilibria work in the NM-Co-Fe system. As mentioned in the last section, only compounds of the type AB, and A2B,7 are of importance for permanent magnet applications. However, it was reported (46,47,48) that optimum properties of sintered RECo, magnets were attained only if the starting material was hyper stoichiometric, i.c., contain a few percent of the RE2Co. phase. So, the study of MM-Co-Fe system was restricted to the composition range between MM2(Co,Fe)7 to MM2(Co,Fe)77.

The phase equilibria work is expected to show the different phases that coexist in the MI-Co-Fe system. Of these only a few may be of importance for permanent magnet was. Curic temperature is an important magnetic characteristic of permanent magnetic materials. Takes, to get a first hand information on which alloys have favourable magnetic characteristics, it was decided to determine the Curic temperatures of various RE-Co-Fe alloys. Since it is known that the Fe addition affects the magnetic easy axis of the A2B17 type phase, it was also felt that the easy magnetization axis of the known and the new phases in the MM-Co-Fe system should be determined as a function of Fe content.

In the fabrication of RE-Co acquets, various technical like sintering, liquid phase sintering etc. are used for developing optimum properties. To choose suitable sintering and annealing temperatures for the temperature his-Co-Fe alloys, a knowledge of the melting points of various alloy compositions is very essential. Thermal analysis methods are suitable for this purpose. Hence, a high temperature thermal analysis apparature considered to be set up for determining the melting points of the MM-Co-Fe alloys.

#### CHAPTER II

## THE MAL ATALYSIS APPARATUS

Since in the fabrication of the magnets the ternary alloys of mischmetal, echalt and iron are to be used, a knowledge of their melting points is necessary for choosing suitable sintering temp ratures. Thereal analysis methods are suitable for establi hing the solid-liquid transition temperatures. Hence, a thermal analysis facility was set up.

# II.1 Design and Description of Apparatus:

during thermal analysis, a large amount of sample and slow heating and cooling rates have to be used. In the present investigation of the RE-Co-Fe systems it was expected that only small amounts of materials would be available for thermal analysis. Hence, a thermal analysis apparatus which would be sensitive enough to show reasonably large thermal arrests even with small amounts of materials was required to be built. An already fabricated water-cooled furnace chamber (49) suitable for use with high vacuum as well as with inert gas atmosphere, was available for conversion into a thermal analysis apparatus. As shown in Fig. II.1, the chamber was in three parts - a bottom plate, a central cylindrical section and a top plate. A small furnace, 1 1/4" dis x 12" long, was

• . • For the purpose of thermal analysis of the support of by three for the purpose of thermal analysis of the sample, a Pt-Pt + 10 percent Rh thermosuple was inserved through the top plate.

The furnice had to be designed on the beside of the requirement that if should cork at temperatures in the range of 1000°C to 1400°C (or 1500°C) in inert atmosphere or high vacuum and that it should be possible to get heating and cooling rates as small as desired. The choice of material for the furnace was limited to only a few materials like Ta, Me and Pt-Rh alloys. Of these, only Ta and Mo were tried. Since a small furnice with sufficient electrical resistance was desired, Ta (which has higher specific resistance than Mo) was first chosen even though its use necessitated the use of high vacuum. Soft, annealed 0.02° did Ta wire was relied into a strip of 0.008° thickness and a furnace with 2.5 Ap total resistance was fabricated. The difficulties with this furnace

were that (1) it meeded maintenance of high vacuum (4x10-5 mm Hg or better) for a long time. (2) since the furnace had to be heated slowly, time of operation lengthened and (3) long period in high vacuum caused large volatilisation of material. the Te furnece failed to operate satisfactorily, No winding was The available He wire was of 0.025" dis and was comparatively more stiff than the available Ta wire. Because of this the Me wire could not be relied down to a thin strip of higher resistance. A Mo would furn ce of the same size as the Ta furnace gave a total resistance of only 1 A and the control of heating or cooling rate was found difficult. To increase the electrical registance of the Summee, two concentric No wound furnaces connected in series were tried. Two recrystallized alumine tubes (cylindrical parts of recrystallized alumina crucibles) of 1" 0.D. x  $1\frac{1}{2}$ " long and 1 1/4" 0.D. x 1 1/6" long respectively were used for making the two furnaces. were put concentrically and connected in series using a stainless steel connector. The total resistance of this two-layer furnace was about 3 A.

In the imitial runs at high temperatures (1300°C to 1400°C), the Pt vs Pt + 10 percent Rh thermal analysis thermocouple got stuck to the locally made alumina protection sheath kept in the melt and the thermocouple wires snapped during its removal from the sheath. To avoid this, it was

necessary to keep the thermocouple in a tube which could be kept in the aluming sheath so that no damage is done to the theremocounts. Thin walled pure all into table of 1/8" would have been the ideal choice. But, since it was not available, a t in walled (0.01" wall thicknoon) steinless steel tabe of 1/8" dia, closed at one and by fusing it, was used. The use of stainless steel tube, however, routrieted the operation of the further upto 1400°C. The ther occupie was inpulated from the stainless steel tube by putting thin Al203 sleeves on the wires and by keepin, the het junction in alumina powder put at the bottom of the shairless stoom tube. To thin stainless stool tube, if expessed to atmosphere at high tempcratures, was expected to oxidise and fail. So, it was connected to a gas head made of brass for flowing argon gas. Purified argon gas was inserted into the stainless steel tube through a syringe needle. The stainless steel tube-gas head was supported by a holder as shown in Fig. II.1.

its operation. Argon gas purified by passing of over hot copper turnings and titanium chips was used for this purpose. The gas was inserted through the bottom plate into the chamber and let out from the side of the furnace chamber through an oil bubbler. For ensuring that all the air was replaced by argon gas before heating the furnace, a vacuum pump was connected to the system as shown in Fig. II.2. The furnace chamber

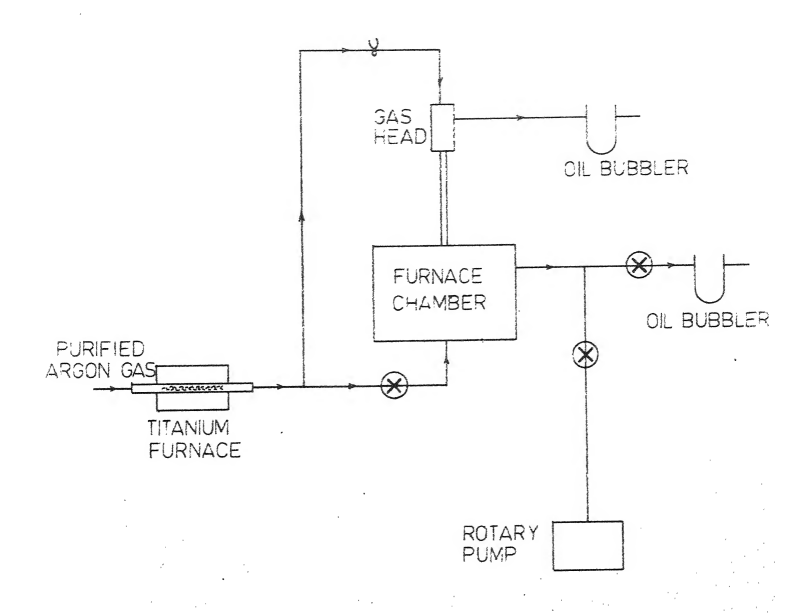


FIG. II. 2. BLOCK DIAGRAM REPRESENTING THERMAL ANALYSIS SET-UP

was evacuated to about 20µ Hg and argon was flushed through the system 2 or 3 times before commencing heating of the furnice.

Since small amount of material was to be used for thornal analysis, the total amount of heat liberated or absorbed at the transformation point was expected to be small. The sucl! heat effect was expected to make the thermal arrest of short duration and with relatively large heating or cooling rate (eg. 2°C/min) if right not be possible to see clearly the small thormal arrest. Because of this, very small heating or cooling rate ( \sigma 0.50 C/min) was desired. The total resistance of the furnace being small, a very fine control of voltage was necessary to obtain slow heating and cooling rates. This was achieved through an arrangement shown in Fig. II.3. A stabilized stepped down voltage was fed to three Dimmerstats set in tanden, the first the were used to step down the input voltage suitably so as to get fine voltage central over the whole range of the Dimmerstat and the third one was used to increase or decrease the set voltage by small amounts.

For thermal analysis, the output of the thermocouple was fed to a recorder through a celd junction (molting point of ice) so that the changing mv, corresponding to the changing temperature, with time could be recorded directly. The mv generated by the thermocouple in the useful temperature range was between 8 to 14 mv. So, one could directly use a

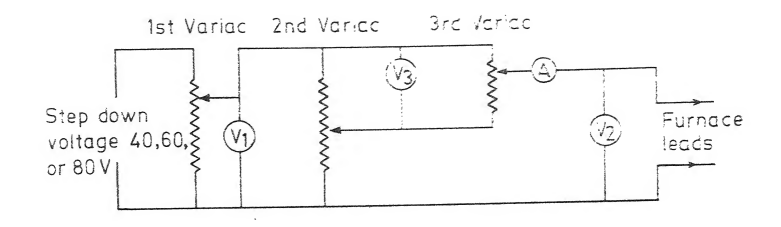


FIG. II.3. FURNACE POWER SUPPLY CIRCUIT

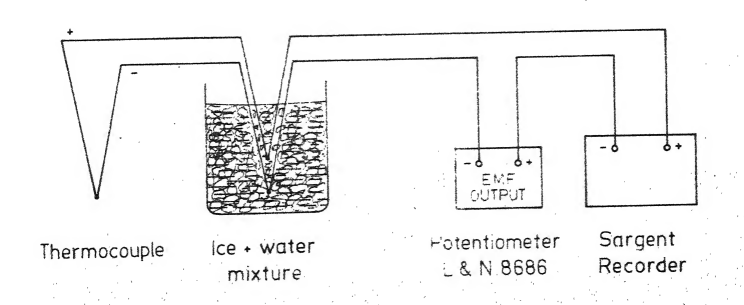


FIG. II 4 THERMOCOUPLE OUTPUT MEASUREMENT CIRCUIT

recorder with full scale deflection of 10 or 20 my and record the thornel arrest. However, to large recorder range required (0-10 or 0-20 my full scale) ande the measurement rather inaccurate because the changing my per unit time was rather low ( 0.01 mv/rin) at the transformation point. Hence, it was thought to make the measuring circuit more sensitive. Since the major part of the my reading was useless for our purpose, it was thought that if an end of known value was fed against the thermocouple enf and was kept constant throughout the experiment, them a more sensitive scale such as 0-1 my range could be used. A Letds and Northrap, Model 8686 potentiometer was used through which a predetermined omf could be fed against the thermoccupic end. The accuracy of the petentiometer is ± 0.007 mv and hence, a very fine control of emf was possible. The unbalanced emf of this system was fed to a Sargent recorder (Fig. II.4) with a 10" chart span and with 0-1 my full scale defloction.

The RE-Co alloys are reported to react with  ${\rm Al}_2{\rm O}_3$  crucibles (50). Commercially available  ${\rm Al}_2{\rm O}_3$  crucibles, which are usually sintered at low temperatures ( ${\rm in}$  1400°C), are very perous and cannot be used. But, since no other suitable crucible material was available, it was considered to use  ${\rm Al}_2{\rm O}_3$  crucibles sintered at higher temperatures. The facilities available in the Institute were used for this purpose. 99.9 percent pure  ${\rm Al}_2{\rm O}_3$  powder ( ${\rm in}$  5µm in size) was blended with

water and for optimum plasticity of the reterial, the blend was maintained at a pH of 3-4 using HCl. Then the cruciblus and sheaths of required sizes were node by slip casting the blended alumina in plaster of paris moulds. An allowance of about 30 percent was given for shrinking during sintering. The cast crucibles and sheaths were presintered at 1000°C for 10 hrs. After machining, if necessary, the presintered crucibles final sintering was done at 1700°C for 2 hrs in a zirconia pot furnace. Crucibles of high density and wall thickness of 1/16" could be propured by this sintering operation.

#### II.2 Tosting and Calibration:

temperature characteristics of the thermal analysis set up and to determine how slow heatin; and cooling rates could be achieved with the two-layer furnace assembly. The power supply to be furnace was raised in steps of 2 volts and the furnace was allowed to stabilise its temperature before each successive increase of input voltage. The temperatures attained corresponding to different voltages are listed in a Table in Appendix I. In the lew temperature region (upto about 800°C), the furnace usually took about 8-12 minutes to stabilize its temperature after every increment in voltage (2 volts) whereas in the high temperature region (> 800°C)

the that to stabilize temperature was about 4-8 mts. Or to whole, the temperature of the furnece was raised at the rate of 3-400/min. After attailing the desired temperature the desimed cooling or heating rates were obtained by slightly reising or lowering the applied veltage to the furnace. The histing and conling notes could be controlled effectively by this nothed. For example, a change of the veltage by about 0.6 volts at 1000°C productd a hortin; rate of about 1°C/min. Si ulerly, for obtaining a cooling rate of 100/min around the same temperature, a voltage change of about 0.5 volts was meeded. For slower heating or cooling rates, smaller voltage changes 0.2-0.4 velts were needed. A chance of voltage by about 0.3 volts produced a heating rate of 0.500/min and Su same cooling rate needed a change of input volume by about 0.25 volts. The slowest heating or choling rate achieved was about 0.40C/min. Still slower heating or cocling rates could not be achieved for the following reasons: (1) when Voltage lower than 57 was impressed or the third Dimmerstat, the heating or cooling rates could not be centrelled, (2) since voltmaters of smaller margin (0-1 V) were not available, the voltage changes could not be measured accurately. At higher torperatures (1500°C an' roru) central of heating and cooling rates was relatively easy because of the increase in furnace resistance.

After initial testing of the funcce assembly, a test run with one of the calibrating specimens (Germanium) was made. 15 grs of 99.9 percent pure germanium, obtained from Semi Elements Inc., N.Y., U.S.A., was kept in a 1 I.D.x1" long Algo, cricuble supplied by Headway Coramics, Varanasi. A small fused silica sheath was used. for protecting the thorrocomple fr. a coming in contact with the material. It was inserted into the material and then two crucible was kept in the furnace. The furnace cumber was assembled and the thermocouple was inserted through the top plate. The system was evacuated to about 20 Hg pressure and flush d wit' purified argon gas 3,4 times. The chember was finally filled with purified argon gas. During the run, publified argon gas was allowed to flow through the chamber as well as the thereocouple assembly and bubbled through the oil bubblers at a rate of 4 to 5 bubbles/min and 1 bubble/min respectively. Since about 45 volts supply was needed to raice too furnace temperature to about 960°C (see Appendix I), a stabilized stopped down voltage of 60V was fed to the first Dirmerstat and the second Dimmorstat (with 45V input veltage) was then centinuously monitored to raise the temperature of furnace at a rate of 4-5°C/min. The third Dimmerstat was set at zero during the initial heating up of the furnace. It was ensured that the furnace chamber was properly cooled during heating

up of the furnace. Ice was put in the thormoccupie cold junction both. The strip chart recorder was calibrated and connected to the thermocouple omf measuring circuit. The furnace temperature was raised to about 930°C using the second Dirmorstat. Then the differential voltage ( \$50) from the third Dimmerstat was used for further clow heating and cooling. In the first run, a heating rate of 2°C/min was used and the output em was plotted on the recorder. A small thermal arrest was seen round 8.85 my cost output (corresponding to about 940°C). So, the furneec was allowed to rice to only 950°C. Then it was crold slowly at a rate of 1.5°C/ min. This time, a comparatively large thermal arrest was found at 8.86 mv. To wike the thereal arrest still larger, shower heating and cooling rates (1°C/min. 0.5°C/min and 0.4°C/min) were used and the time temperature curves were traced on the recorder. A small variation in the emf of the arrest point, ± 0.015 my around the mean value, 8.85 my, was found in different heating and cooling cycles. A cooling curve obtained with 0.4°C/min cocling rate is shown in Fig. II.5(a). After the thermal analysis muns were made, before shutting off the furnece power, the valves near the oil-bubblers were closed. This was done to provent sucking up of bil into the furnace chamber or into the thermoccuple assembly during cooling of the furnace. Then, the furnace was allowed to each by putting

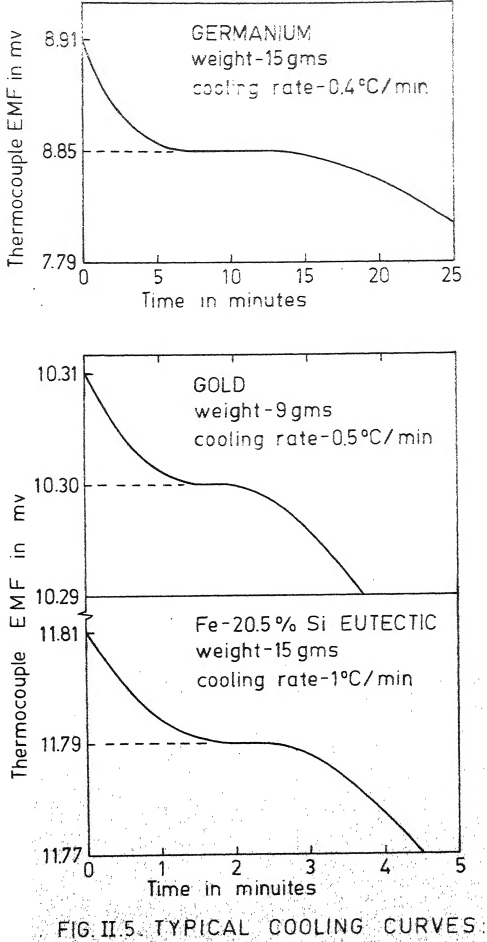


FIG. II.5. TYPICAL COOLING CURVES

(a) Germanium (b) Gold (c) Fe-Si eutectic

off the power supply.

Calibration is usually made against the malting points of pure metals. For calibrating the thereoccupie in the temperature range of interest (1000°C-1400°C), germonium and gold having molting points 937.4°C and 1063°C were used. The third colibration was planned with silicen beging a nelting point of 1/1000. But, due to the use of scripless steen thermocouple sheath, milicon could not be used as a calibration retarial because the multing point of steinless steel (  $\simeq 1420^{\circ}$ C) is very close to the molting point of silicon. Since no other non-reactive pure metal with accurately known molting point in the temperature range of 1100°C and 1400°C was available, it became recessary to use an entectic allow of known cutoctic point. Fe-20.5 wt. percent Si cutoctic was found suitable for the present work. There is some uncertainty in the reported cutectic temperatures of Fe-Si ( 20 wt porcent) alloy. Houghton and Book - (51), using high purity materials, reported an outectic temperature of 1195°C for Fe-20 wt. percent Si. Murakani reported a temperature of 1205°C for Fe-23 wt percent Si (52). Fhragmen (53) and Obawa et al. (54) reported an eutectic temperature of 1200°C for the Fe-21.2 wt percent Si and Fe-22 wt percent Si alloys respectively. On the basis of these data Hansen and Anderko (55) has shown the entectic temperature as 1200°C.

The same temperature with a possible uncertainty of ±5°C has been used in this calibration as the entectic temperature of Fe-20.5 wt percent Si. About 15 gum of the entectic alley was prepared by are melting electrolytic iron flakes obtained from Gallard Scholsinger Hfg., Co., F.Y., U.S.A. and silicon obtained from Semi Elements Inc., N.Y. U.S.A.

In the case of Co and the Fo-Si allow, the arrest points were found to drift comowhat and hance mean values of thornal arrest end of all the beating and cooling runs were used for thermocouple colibration. For gold, no variation in the arrest point was observed (Table II.1). With the help of the known solting temporatures and the thermocouple enf values corresponding to the thermal arrests, a colibration curve, Fig. II.6, was drawn. Since there was some drift observed in the thermal arrests of Ge and Fu-Si all y and because there was a small uncertainty in the melting point of Fo-Si alloy (uncortainty limits are should by horizontal and vertical bars in Fig. II.6) the temporature measured with the help of the thereoccupie will have a small uncertainty. It is, however, clear from the calibration curve that this uncertainty is small, about ±5°C. The therrocouple was calibrated only in the temperature range 930°C to 1200°C. However, from the amosthness of the calibration curve, it is expected that it can be extended safely to about 1350°C or 1400°C.

TANTE II.1
Thermocomple Calibration Data

Serial No.	Materi ]	Thermocouple MF corresponding to the molting point of the moterial	Melting Point
1	Gornyainn	8.85 <u>+</u> 0.0 <u>1</u> 5 : v	93 <b>7.</b> 4 <sup>0</sup> 0
2	Gold	10.3 m	1063°G
3	Fc-20.5 wt per-	11.785±0.025 =v	1200° <u>±</u> 5°0
	cent Si eutectic		

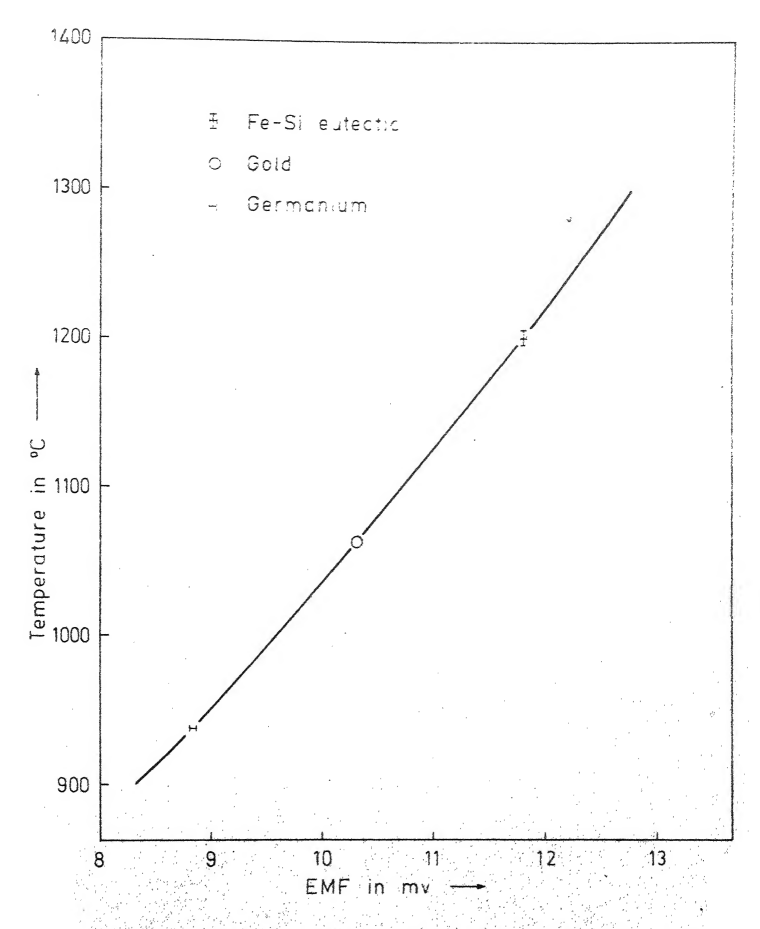


FIG. II.6. THERMOCOUPLE CALIBRATION CURVE

# CHAPTER III EXPARIA LEAL PROCEDURE

The present investigation involved the determination of

- 1. Phase equilibria in RE-Co-Fe system
- 2. Solidur and liquidur temperatures of the alleys (Thornel Arrivais)
- 3. Curi: Temperatures

and 4. Easy axis of magnetization

#### III.1 Phase Equilibria in RE-Co-Fc System:

For establishing the phase equilibric in RE-Co-Fe system, the experimental procedure followed consisted of the following steps: a) Melting of alloys, b) A nealing,

c) Metallographic analysis, and d) X-ray diffraction analysis.
Melting:

Materials Preparation: The alloys were prepared by using 1) Mischmetal supplied by Mischmetal and Flints Pvt. Ltd., Alloppey, Kerala, India, containing 85-90 wt percent Rare Earth (RE), 6-10 wt percent Iron (Fe) and 1-4 wt percent impurities and the rare earth composition was Co: 45-50 wt percent; La: 20-25 wt percent; Nd: 15-20 wt percent; Pr: 4-6 wt percent; Sa: 1-4 wt percent; Y and others: 1-2 wt percent,

ii) Cobalt (9929 percent) supplied by Seri Blements Inc.,
N.Y., U.S.A. and iii) Blectrolytic iron flates (99.9 percent)
supplied by Gallard Scheldinger Mfg. Co., N.Y., U.S.A. Since
the exact amounts of individual rare eart' of matrix in mischmotal were not known, an average composition was assumed
busing on the supplier's analysis give, above. The average
composition used for calculating the rare carts nolocular
weight in mischmotal (MM) was 90 wt percent total RB; 8 wt
percent Fe and 2 wt percent impurities and the individual rare
earth compositions were 49.21 wt percent Ce, 23.31 wt percent
La, 1813 wt percent Nd, 5.18 wt percent Pr, 2.59 wt percent
Sh and 1.55 wt percent Y. The calculated rare carth molecular
weight on this basis was 140.11 gms. This polecular weight
of RE was used in all calculations of the amounts of RE to
be taken for different alloy compositions.

The mischnetal as supplied was in the form of biscuits conted with grease to protect it from atmospheric exidation. Before cutting the mischmetal to pieces of required size, the grease was removed using acctana. The cut pieces were then ground to remove any exide layer that was present. The pieces were weighed according to the alloy composition so as to make the total amount of alloy between 25 and 30 grs and the weighed pieces were kept in acctone till they were used for molting. Cobalt was in the form of a cast butten

and it was crushed to pieces of required size using a harmor, Iron was in the form of flokes; they were ground to remove the traces of rust. The amounts of iron and cobelt required for a given melt were weighed out and kept separately till required for melting.

Molting Furnace: The factors to be considered in a molting operation are: making points of the elements, amount of material to be melted, reactivity of the components, contamination from crucible metarial. The menting points of the three components in RE-Co-Fe allows are In: Fo: 1539°C and Co: 1492°C. Out of these times notels, mischmotal is very reactive and is reported to react with even alumina crucibles (50). So, it was thought proper to carryout the molting under protective argon atmosphere and without using a refractory crucible. A water cooled copper hearth ere furnace with men-compunable tangston electrode was co-sidered to be the best melting furnece. An available D.C. are furneed was thus node use of for molting the RE-Co-Fo alloys. The cracible and the electrode were elelosed in a glass chamber which could be evacuated and an inert gas atmosphere could be unintrained in it. The electrode was fitted to the top netal part of the glass chamber through O-ring joints so that the electrode could be raised or lowered with the help of a serow. This was necessary to adjust

the grap between the electrice and the charge (usually kept between 1/8" to 1/4") so that are could be struck without trucking the charge with the electrode. It conside clean, contamination free colling. A D.C. are welder (1000 amps, 40 V) was used as the are power supply. The different centrals - start and weld centrals - varying the welding current from 300 amps at 0 to 1000 amps at 100 or the central dish were available. With the help of these centrals the are pressure could be centralled. An automotic timer (0-180 sees) which could be set according to the required duration of colting was also available in the welder start circuit as as to shut off the are current after a pre-selected time.

Gas Purification System: The common bottled argo: gas contains moisture and oxygen both of which are detrimental to the Rare Borth alloys. Hence, it had to be forther purified before putting it into the are furnace charber. For this purpose, a gas purification system was constructured. As shown in the block diagram (Fig. III.1), it consisted of several GaCl<sub>2</sub> and P<sub>2</sub>O<sub>5</sub> absorber tubes and furnaces with Cu turnings and Ti chips. The Cu turnings and the Ti chips were kept in four ½" dia x 30" long quartz tubes over a length of 6".

The quartz tubes, in turn, were kept in a 2" dia x 15" long michrone-wound ference. The furnace was kept at 700°C so that the titerium remained in the α state (because it has higher

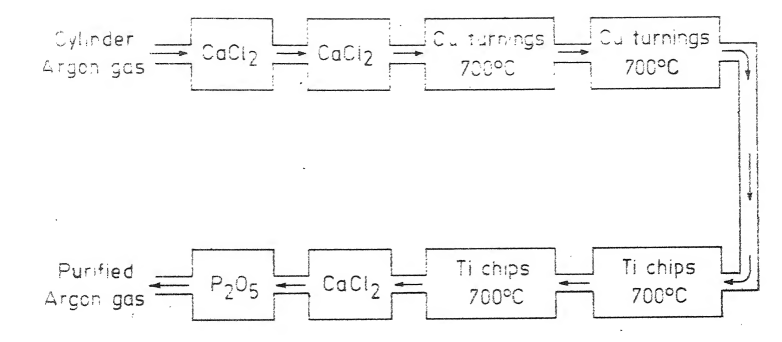


FIG. III 1 BLOCK DIAGRAM REPRESENTING GAS PURIFICATION SYSTEM.

solubility of  $0_2$ ), at the same time the reaction rate between oxygen and bu turnings or Elchips accelerated. Argon gas from the gas cylinder was bassed through the CaCl<sub>2</sub> tubes to remove the bulk of the moisture and then through hot Cu turnings to relove  $0_2$  and traces of  $0_2$ . The gas was then passed through Ti chips to remove  $0_2$ ,  $0_2$  and  $0_2$ . The gas was finally passed through CaCl<sub>2</sub> and  $0_2$ 0, to remove the last traces of moisture. This gas was used for flushing and final filling up of the are furnace chamber.

Mediting Procedure: The weighed materials were always charged in the copper crucible in a particular sequence so as to keep the evaporation losses the same and at the minimum. Mischintal. which is the lowest mediting component (n.p. an 900°C) amongst the three components, M., Co and Fe, and has reasonable vapour pressure at 1 atm. pressure, was always kept at the bottom and was covered completely by iron and cobalt pieces. Thus, initially, the are was always struck between the tangsten electrode and an iron piece kept at the top. Before each mediting operation the furnace chamber was evacuated to about 20µ Hg pressure and flushed with purified argon gas from the gas purification system. This procedure was repeated 3 or 4 times to ensure removal of all the air from the furnace chamber. Finally, the furnace chamber was filled with purified argon gas. In the first melting, by setting the start and weld

controls at 12 and 10 respectively, lover arc pressure was used to avoid sputtering of small pieces of reterial. The time was set at 20 secs to have a short meltin, tile. But, in the subsequent meltings of the alloy button the start and weld controls were set at 18 and 15 respectively and the timer was set at 30 secs so that higher are pressure and longer melting times could be used for proper hopogenization of the material. Each allow was melted 3 to 4 times and at each remelting the maloy button was put upside down so as to produce better homogenization. After melting, the alley was weighed to find out the total loss. It was found that the next mum loss was less than I percent. Some of the alloys were chemically analysed for Fe and Co to check whother or not the alloys with compositions close to the intended one could be prepared.

For phase analysis of the allows, the alloys had to be properly homogenized before they were subjected to x-ray diffraction and metallographic analysis. Since the alloys were rather reactive and each netrod of analysis was spread over several days it was necessary protect them from exidation. For storage of the alloys till they were needed a vacuum desiccator kept at 750µ Hg pressure was used.

Amnealing: The as cast samples were broken into pieces of suitable size and the surfaces of the pieces were ground to remove any oxide that might have formed during melting or sto-The alloys were sealed in evacuated quartz tubes at a pressure of 20 x 10<sup>-3</sup> rm Hg. Sufficient space was loft between the walls of the quartz tubes and the samples to avoid breakings of quarts tubod due to thermal expansion of the alloy pieces. Long pieces of an alloy were put, wherever possible, because the small pieces had a tendency to get stack (muchamical jasming) in the quartz tubes leading to their breakage when heated. In a betch, five to six such quartz tubes containing different alloy samples were annealed and proper care was taken to identify them. In the beginning, the arc melted alloys were heat treated at a reasonably low temporature to see if any problem, like reaction with quartz tubes, oxidation in the residual air etc was encountered during annualing. Initial annealing was done in a vertical tube resistance furmace at 500°+1°C for 5 days. The alloys were quenched in tap water. No reaction between the quartz tubos and the alloys could be detected. Some of the alloys after their low temperature ennealing showed three phases while the same samples showed only two phases after annualing at 900°C for 5 days indicating that the 5 day annualing at 500°C was not sufficient for homogenization. Subsequently, it was found that a 4 day amoral at 900°C (or higher time amoral at 900°C) produced the same microstructure of the alloys as that in a 5 day or a longer anneal. Hence, all alloys were amorated at 900°C for 4 days and finally quenched in top water. The annealed alloys were also stored in the evacuated desiceator.

Metallographic Analysis: A small and representative portion of the annualed alloy was taken from the annualed sample for motallographic analysis. A vertical section from the nelted button, which would show up any segregation or inhomogeneity of the phases, was generally solucted for this purpose. The samples for metallography were mounted in Jucite. The mounted samples were polished on 1/0 through 4/0 enery papers. Finally, the samples were polished on a wheel covered with microcloth using 0.5u alumina powder suspended in water as the abrasive. After polishing, the samples were thoroughly washed with water, rinsed with alcohol and dried under a high speed blower. The sampler were checked under an optical microscope in the ac-pelished condition to detect the presence of inhomogeneitics and oxides. The polished samples were then etched to reveal the microstructures. The etching reasont used (Nital) consisted of 2.5 %1. conc. ENO, in 97.5 ml of pure methanol. All the samples were exclud to the same extent so that different samples processed in the same manner could . be compared. After etching, the samples were thoroughly washed With water, rinsed with alcohol and dried under a blower. The concentration of the acid in the etchent and the etching time were varied for some of the samples to obtain better contrast among the places. All percent mital solution was found suitable for revealing three phases in some of the 3- phase alloys.

X-ray Diffraction Analysis: About 3 to 4 gms of the annualed sample was used for x-ray diffraction work. Most of the alloys were very brittly and so it was easy to prepare -300 mash powder using steel and agate mortars and postles. But, some of the high Fc alloys were found to be comparatively tough and grinding had to be continued till the whole material passed through the sieve. Since the diffraction patterns with the as-crushed powder produced sharp diffraction lines, the powders were not amnealed before taking diffraction patterns.

For phase identification, the diffractometer patterns were obtained using a G.E. XRD-6 diffractometer with unfiltered Cr-radiation at 30 KV, 17 mA. The criterion used in choosing the diffractometer in comparision with the Debye-Scherrer technique and the radiation were: high resolution of the diffractometer, high resolution due to the long wave length of Cr-radiation and shorter time to obtain a diffraction pattern. All the diffraction patterns were obtained in the angular range of 30° < 20 < 145°. The diffractometer conditions used in

recording the diffraction potterns were scanning speed of  $2^{\circ}/\sin$ , slit system  $3^{\circ}$  MR | MR / 0.2°, time constant of 4.0, a range of 500 cps and a chart speed of 30 //r. Unfiltered radiction was used so that the weak peaks could be detected. Some of the very weak peaks, however, could not be clearly scen in the diffractometer traces because of reasonably large noise in the recording system. For phase analysis the diffraction traces of pure or nearly pure phoses were used as standard patterns for comperison with other diffraction traces. The diffraction patterns of pure or nearly pure phases were taken using a 114.6 rm die Debye-Scherrer Castera. These diffraction patterns were used for proper identification, through indexing of the diffraction patterns, of phases. An exposure of 9 hrs. with unfiltered Cr-radiation at 30 KV and 15 DA gave good diffraction patterns. After the x-ray work was completed, the powdered samples were stored in evacuated and sealed pyrox glass tubes for future use.

# III.2 Thormal Analysis:

For determining the molting points of RE-Co-Fe alloys, the thormal analysis apparatus described in Chapter II was used. Since the RE-Co alloys are reported to react with Al<sub>2</sub>0<sub>3</sub> crucibles (50) special precautions were taken to avoid damage of the furnace assembly due to seepage of material through the crucible. Two concentric, sintered Al<sub>2</sub>0<sub>3</sub> crucibles,

instead of one, were used for keeping the motorial. Since the molting points of the investigated termory alloys could not be guessed from the binary systems, for the first tried run an alloy with the lowest Fe was chosen. It was expected that its molting point would not be far from those shown by the bimary diagrams. An available RE-Co-Fe two-phase (convaining RECo5 and RE2Co17 type phases) allow with least amount of Fe was chosen and it was kept in the locally made  ${\rm Al}_2{\rm O}_3$ crucibles. After keeping the crucibles in the farmece, the furnice chamber was evacuited and flushed with argen gas 3 or 4 times. Since the thereocouple was to be protected with a thin Al203 sheath and it was not known how this thermocouple sheath would behave in the molten RE alloy, in the first run it was kept outside the melt. The funce was heated up slouly and when the temperature reached about 1350°C (moun from the applied voltage, see Appendix I) the thereocouple along with its protection sheath was pushed into the melt so as to start thermal analysis. But, this attempt was unsuccessful because the material in the crucible was found to be in partially rolten condition. Later, a relatively pure MI(Co,Fe)5 alloy close to the bin ry (which was expected to have lower molting temperature) was used for thermal analysis. This time, the thermocouple along with the Along sheath was kept incide the material from the beginning. Thermal analysis run was made following the procedure given in Chapter II.

#### III.3 Curic Temperature Measurement:

For determining the Curic temperatures of RE-Co-Fe alloys, an apparatus fabricated by R.C. Nittal (57) was used. It essentially consisted of a small nichrome-wound furnace kept in a vacuum chamber - a pyrex glass bell jar with a long tube at its top. A sealed end brase tube inserted into the furnace chamber through an 'O' ring joint could be moved up or down for lowering or raising the sample. The sample was kept in a quartz tube hung from the bottom end of the brass tube. For measuring the temperature, a thermocouple was inserted into the material through the brass tube; an alumina sheath was used for protecting the thermocouple. An induction coil kept over the long tube portion of the elember was used for determining the flux change as a function of temperature. A differential method was adopted to measure small changes in induced voltage. A rectifier circuit with provision for continuous balancing of the rectified veltage to a near null voltage was used for converting the A.C. signal to a D.C. signal. The D.C. output voltage from the rectifier circuit was amplified usin a D.C. microvolt amplifier with various D.C. ranges (0-100uV full scale to 0-1000 V full scale) and recorded on a Sargent Strip chart recorder.

recorder. Sufficient care was taken to prevent the vacuum chamber from gettin, heated up.

# III.4 Magnetic Easy Axis Determination:

As has been pointed out in the introduction section, the easy axis of magnetization of RECo<sub>5</sub> and RE<sub>2</sub>Co<sub>17</sub> alloys are either along the crystallographic c-axis or in the basal plane. If the powdered samples consist of single domain particles, then in a small magnetic field the loose powders are expected to align themselves along the east axis. In such oriented samples, the particles will arrange themselves with their easy axis along the direction of the magnetic field but will have random orientations around this easy axis. This aligned speciagm is thus similar to a sample with wire texture. Hence, the oriented specimen diffraction pattern should show similar streaky pattern as is obtained with drawn wires or fibres. Thus the easy magnetisation axis can be determined in the same way as the wire axis is determined.

For preparing the oriented sample of alloy 51, a small amount of -300 mesh powder was taken on a card and a small SmCo<sub>5</sub> magnet was placed below the card. A little juggling of the magnet produced short thin magnetically aligned needles standing erect on the card. These needles were frozen in by spraying krylon clear plastic spray on them. The magnet was then removed and the needles were removed by scraping the paper with a sharp razor blade. Of the large number of needles produced, the thinnest and the most

uniform one (determined by visual observation) was mounted in a 57.3 rm dia Woissenberg Camera and a powder pattern was The diffraction pattern obtained showed basically continuous Debye rings with indications that some of the diffraction lines have non-uniform intensity distribution. Since the allow 51 was somewhat tough compared to some other allow. of higher RE content, it was thought that the surface layers of the particles might have got some plastic flow during the mochanical grinding operation causing apparent randorness and hence were producing continuous Debye rings. To remove the disturbed layers from the particle surface, the alloy powder was subsequently otched by 5 percent nital for 30 secs. diffraction pattern obtained with the etched powder showed clear streaky pattern as is expected from a well aligned specimen. After locating the ideal position of the diffraction spots on the intensity streaks of the diffraction pattern, the position of the same spots on a flat file, kept at the same distance from the specimen as the cylindrical film, was deternined by geometrical principles. Then the standard procedure for wire texture determination was adopted.

#### CHAPTER IV

#### RESULTS AND DISCUSSION

## IV.1 Phase Equilibria:

Some of the rare earth elements present in Mischmetal have complete solid solubility whereas others have extended solubility among themselves (33). For an individual transition metal (Mn, Fe, Co or Ni), the binary systems With different rare earths exhibit similar binary phase equilibria. This is in general true also for the nontransition metal-rare earth binary systems. Hence, when rare earth elements are present together, as in Mischmetal, the phase equilibria is expected to be similar to that of the rare earth binary systems. The work of Fishman and Crowe (34) on the MM-Mg system indicate that this assumption is a reasonably good one. Assuming that the above observations are in general true, the MM-Co-Fe system can be represented as RE-Go-Fe system, where RE stands for the rare earth elements The RE-Co-Fe system was studied in this inpresent in MM. vestigation through metallographic as well as x-ray diffraction techniques. For phase analysis by x-ray diffraction, standard patterns of the  ${\rm RECo}_5$  and  ${\rm RE}_2{\rm Co}_{1.7}$  phases (57) were utilized. The phase analysis results of the 900°C annealed alloys are tabulated in Table IV.1. The isothermal section of the RE-Co-Fe system drawn on the basis of the above analysis is represented in Fig. IV.1. Some of the alloys were

TABLE IV.1
Phase Analysis of the RE-Co-Fe System

Alloy			sition of	75 1	Phase Analysis
No.	RE RE	in at. po		Metallo- graphy	X-ray Diffraction
	T.L.	Co	Fe	3	
	•				
8	16.66	63.34	20.0	3	RE <sub>2</sub> Co <sub>7</sub> +RECo <sub>5</sub> +S
12	20.0	65.0	15.0	2	RE <sub>2</sub> Co <sub>7</sub> +RECo <sub>5</sub>
14	20.0	60.0	20.0	3	RE <sub>2</sub> Co <sub>7</sub> +RECo <sub>5</sub> +S
15	16.0	64.0	20.0	3	RE <sub>2</sub> Co <sub>7</sub> +RECo <sub>5</sub> +S
16	15.0	60.0	25.0	2	RE <sub>2</sub> Co <sub>7</sub> +S
17	15.0	55.0	30.0	2	RE <sub>2</sub> Co <sub>7</sub> +S+A
18	15.0	50.0	35.0	2	S+A
19	15.0	45.0	40.0	2	S+A
22	17.0	43.0	40.0	2	S+A
23	13.5	46.5	40.0	2	S+A
25	14.0	56.0	30.0	2	RE <sub>2</sub> Co <sub>7</sub> +S+A
28	12.75	47.25	40.0	2	STA
30	15.0	65.0	20.0	3	RE <sub>2</sub> Co <sub>7</sub> +RECo <sub>5</sub> +S
31	11.75	38.25	50.0	2	S+X
33	18.0	35.5	46.5	2	A+Y
34	17.5	32.5	50.0	2	A+Y
35	11.0	34.0	55.0	2	S+T+X
39	13.0	52.0	<b>35</b> • 0	2	Sta
41	12.11	62.33	25.54	2	S+A
42	18.0	42.0	40.0	2	Sta
43	16.5	33.5	50.0	2	S+A
47	12.5	52.5	35.0	2	S+A
48	13.5	61.5	25.0	2	S+ù
50	12.75	62.25	25.0	2	S+A
51	. 15•0	53.0	35.0	2	S+A
52	19.0	51.0	30.0	2	RE2 <sup>C</sup> O7+A
53	16.5	28.5	55.0	2	T+Z

churically analysed. The intended and a slysed compositions shown in Table IV.2 indicate that the losses are small. Since the analysed compositions are very close to the intended ones, the phase equilibria shown in Fig. IV.1 has been drawn or the basis of the intended compositions of elloys.

In Figure IV.1 several dash dot links are draw, slow certain constant RE contents to represent the hypothetical  $\mathbb{A}_{\mathbf{x}}^{\mathrm{B}}$  quesi-bin ries corresponding to the high cobact RE-Co phoses Of those quasi-binary lines, the AB2 and A2B17 lines are drawn between the RE-Co and RE-Fo binorial because in both these system the AB2 and A2B17 type plases exist and extension of those phases from one binary to the other as exist along these lines. On the other hand, the other  $\mathbb{A}_{\mathbf{x}}\mathbb{B}_{v}$ lines are drawn only upto the middle of the terrary diagram (those  ${ t A_{f x} B_{f y}}$  compounds are not found in the RE-Fe system) to indicate that these  $A_{\mathbf{x}}B_{\mathbf{y}}$  phases may extend to some extent along these lines. In this investigation only a part of the RE-Co-Fe system was studied. In the investigated region of the RE-Co-Fe system, besides the A2B17 type phase two new phases, A-phase and S-phase, were found to exist. A ong the investigated alloys, only two alloys, alloy 12 with 15 at. pet. Fe and alloy 14 with 20 at. pet. Fo, were found to have the RE2Co7 type phase as the rajor phase; the alloy 14 was found to have three phases (Fig. IV.2) whereas the alley 12 was

TABLE IV.2

Chemical Analysis of Several RE-Co-Fe Alloys

Alloy No.	Trte	Compositi ended	Composition wt percent		
managana daganagan katalangan sagangan sa	Fe	Go	Fe	Go	
a.		•		:	
30	15.84	54.34	15.3	54•€	
35	46.42	30.29	46.2	30.1	
41	20,99	54•04	20.8	53•4	

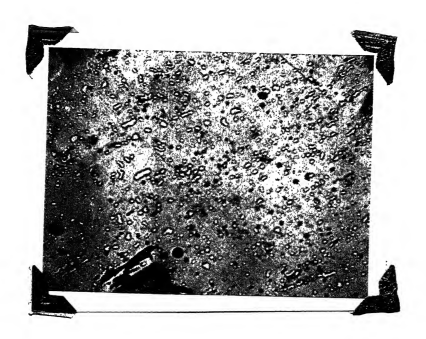


Fig. IV.2 Microstructure of Alloy 14 (200 X)

Photograph shows only two phases; RE<sub>2</sub>Co<sub>7</sub>-type phase (dark matrix) and RECo<sub>5</sub> type phase (bright second phase)

having only two phases. The second and third phases in these alloys were in very small amounts and hence, these alloys could be considered as boundary alloys. Since the RE2Co7 phase exists in the RE-Co system of Ce, La and Nd, tha three major components of MM, it is expected that the Major, phase exists. The RE2Co7 type place at low Fe concentrations was not studied. From the phase conjysis of the RE-Co-Pe alloys, it appears that the RE2Co7 type phase extends from the RE-Co binary upto at least about 20 at. pet. Fe. One charaeteriatic feature of the extension of the  ${\tt RE_2Co_7}$  phase into the . t rmary is that the phase region deviates from the hypothetical A2B17 line towards higher (Co,Fe) side. While no data on the phase extension of the  $\mathbb{A}_2\mathbb{B}_7$  type phase is available in the literature, some observations have been nede with respect to AB, compounds. Buschow (30) and Taylor and Poldy (58) could not produce single phase samples of  $RE(Co_{1-x}Fe_x)_5$ , where RE = Sm or Y, with larger amounts of Fe (>5 percent) unless the stoichiometry was allowed to shift towards higher (Co,Fc) side.

The S-phase appears to be stable over a reasonably wide Fe concentrations. The lowest and highest Fe contents in the S-phase were estimated through the microstructure and x-ray diffraction analysis. The x-ray diffraction patterns of alloys 8,14,15 and 30 showed diffraction lines due to the

 $\mathrm{RE}_{2}\mathrm{Co}_{7}$ ,  $\mathrm{RECo}_{5}$  and S phases. On netallographic analysis. the same alloys revealed two phases when 2.5 percent Nital was used but on using 1 percent Nital three phases could be clearly seen (Fig. IV.3), Colouration produced on one of the phases masked the third phase when 2.5 percent Nital was The alloys 8 and 14, out of the four three phase alleys. contained the third phase RECo5 in quite small indicating that these alloys were close to one side the three phase triangle. In an attempt to draw three phase triangle, it was found that the three phase corner at the S-phase is near about 21 at. pct. Fe. Thus. the lower Fe limit of the S-phase is possibly at 21 at. pc+. Fe or is at a somewhat lower Fe concentration. At the higher Fe side of the S-phase the alloy 31 has two phases, the amount of second phase being very small (Fig. IV.4). x-ray diffraction pattern of alloy 31 showed the S-phase lines together with a few weak lines possibly due to the second phase. The allow 35 also showed two phases, the second phase was in small amounts and looked the same as the second phase in alloy 31. The diffraction pattern of alloy 35, however, indicated that besides all the lines present in alloy 31, there were some weak lines and that the relative intensity relations of the S phase lines were not quite the same as in alloy 31 or alloy 51 diffraction patterns. indicated that the alloy 35 had three phases - the S phase: the X phase, to the higher (Co, Fe) side of the S phase, and

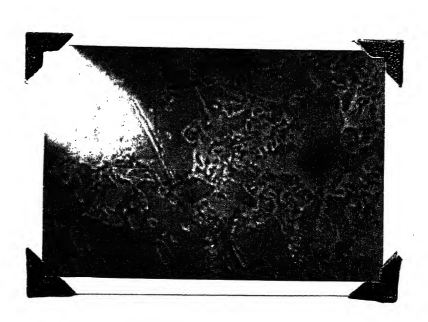


Fig. IV.3 Microstructure of Alloy 15 (1000 X).

Photograph shows three phases; RE2Co7-type phase (dark natrix), E phase (bright second phase) and RECo5 type phase (bright needle like structure) which is in very snall amount.

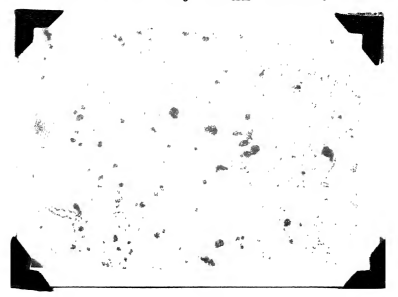


Fig. IV.4 Microstructure of Alloy 31 (200 X)

Photograph shows two phases; S phase (matrix) and
X phase (bright) in small amounts.

the T phase, possibly at a higher Fe content. The x-ray and metallographic observations on alloy 35 indicate that the S phase possibly extends to a Fe content loss than 55 at. pct. Fe. Tentatively, this limit has been shown in Fig. IV.1 as about 51 at. pct. Fe. Metallographic observations on alloy 51 (Fig. IV.5) and alloy 31 indicate that the Sphase region is rather narrow. It is known from the literature that the RE2Co, 7 phase extends from the RE-Co system to the  $^{\mathrm{RE}}2^{\mathrm{Fe}}17$  phase in the RE-Fe binary (59). The S-phase extends almost parallel to the A2B17 line and is very close to it. This indicates that the extension of the RE2Co17 type phase must be to the higher (Co,Fe) side of the A2B17 line. is possible that the x phase is the RE2Co17 type phase. The unidentified diffraction lines in the diffraction patterns of alloy 31 and alloy 35 were too few to identify the x phase as the RE2Co17 type phase. The A phase was found to extend along the same direction of the extension as the RE2Co7 type phase. Alloy 52 showed two phases, almost in equal amounts, in the microstructure (Fig. IV.6) and the diffraction pattern showed the diffraction lines due to the A and RE2007 type phases. The position of allcy 52 relative to the RE2Co7 type phase and the A phase indicates that the A phase region extends at least down to about 35 at. pct. Fe and the RE2Co7 type phase extends at least upto 25 pat. pct. Fe. Alloy 53

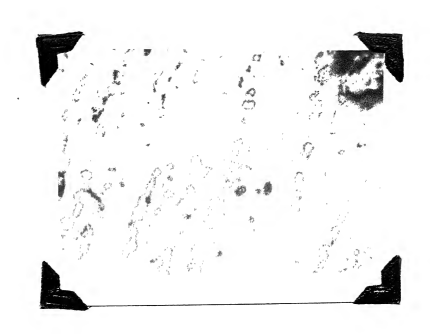


Fig. IV.5 Microstructure of Alloy 51 (400 X)

Photograph shows two phases; S phase (natrix)

and A phase (dark second phase) in small amounts.

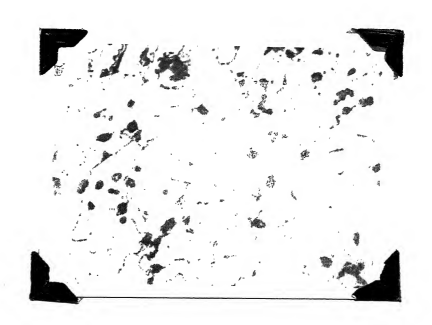


Fig. IV.6 Microstructure of Alloy 52 (400 X)

Photograph shows two phases; RE<sub>2</sub>Co<sub>7</sub> type phase and Λ phase (bright second phase)

at 55 at. pct. Fe was found to have two phases, none of which resembled the A or S phases. The diffraction pattern showed lines which matched with the T phase lines observed in alloy 35. This indicates that the alloy 53 possibly has the T phase and a phase Z phase, with higher RE content then the A phase. Since the alloy 53 has two phases, a three phase region between alloy 53 and A phase is expected. On this basis, the extension of the A phase region to higher Fe side has been shown to be about 53 at. pct. Fe. Alloy 34 showed two phases in the microstructure (Fig. IV.7) and the diffraction pattern showed the diffraction lines due to the A phase and another phase, y phase, possibly a phase with higher RE content than the A phase. Alloy 42 showed two phases in the microstructure (Fig. IV.8) and the diffraction pattern showed the diffraction lines due to the A and S phases. The microstructures of alloys 34 and 42 clearly showed that the second phases in these alloys were different. Thus, alloy 42 and alloy 34 are on either side of the A phoso, A comparison of their compositions indicates that the A phase region is rather narrow. With increase in Fe content, the A phase region was found to extend towards higher (Co, Fe) side and at about 52 at. pct. Fe it crossed the AB5 line. This composition will be useful in determining the crystal structure of the A phase.



Fig. IV.7 Microstructure of Alloy 34 (200 X)

Photograph shows two phases; Bright A phase and dark Y phase. Black spots are probably pits.

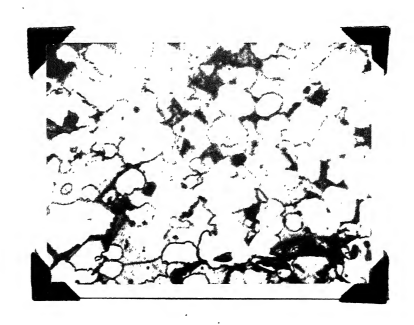


Fig. IV.8 Microstructure of Alloy 42 (400 X)

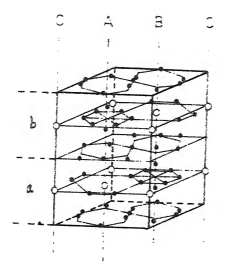
Photograph shows two phases; A phase (dark matrix) and S phase (bright).

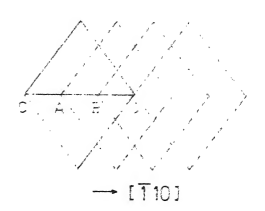
Alloys 17 and 25 were found to have two phases in microstructure but their x-ray diffraction patterns revealed the presence of A, S and RE<sub>2</sub>Co<sub>7</sub> type phases. It is possible that like in alloy 35, the three phases are not revealed by the etching reagent used. On the basis of the x-ray diffraction data these alloys are shown within a three phase triangle drawn between A, S and RE<sub>2</sub>Co<sub>7</sub> phases.

The diffraction patterns of S and A phases are very similar to that of the RE2Co17 phase. For easy comparison, the diffraction patterns of A, S and RE2Co17 phases are shown in Fig. IV.9 in a modified manner. The diffraction peak positions are shown as vertical lines drawn at the appropriate 20 values, the height of the lines represents the relative intensities of various diffraction lines. The diffraction peaks of the A, S and RE2Co17 phases in the low angle region of the diffraction pattern overlap considerably and cause difficulty in identification of the three phases if present together. The diffraction peaks in the high angle region of the diffraction pattern are relatively widely spaced and for phase identification these peaks have been found valuable.

Since the diffraction patterns of the A and S phases resemble that of the  ${
m RE}_2{
m Co}_{17}$  phase, an attempt was made to index these diffraction patterns with the help of the  ${
m RE}_2{
m Co}_{17}$ 

structure. A characteristic feature of the RE2Co17 hexagonal structure (Fig. IV.10) is that it can be thought in terms of stacking of blocks consisting of two atom layers, one with a layer of Co atoms only and the other, a mixed layer of RE and Co atoms. Hexagonal or rhombohedral RE2Co17 structure can be generated by stacking these blocks after shifting each block with respect to the other through a distance 1/3 [1 10] (Fig. IV.11). The sequence ab ab .... generates the hexagonal RE2<sup>Co</sup>17 structure and the sequence abc abc... generates the rhombohedral RE2Co17 structure (Fig. IV.12). One important feature of such layer structures is that the basal plane dimensions, i.e., the a parameters, do not change when the structure changes from hexagonal to rhombohedral symmetry (eg. a parameter of  $\alpha$ - and  $\beta$ -Ce<sub>2</sub>Co<sub>17</sub> are same (15)), only the c parameter changes due to addition of one more block in the rhombohedral structure. The diffraction patterns of the A and S phases being quite similar to that of the rhombohedral RE20017 phase, the structures of the A and S phases may be obtained through proper stacking of blocks, keeping in mind the stoichiometry of the phases. As a preliminary step towards determining the structure of these phases, the diffraction patterns were tried to be indexed. Since many of the diffraction lines of the A and S phases match well with those of the RE2Co17 structure, an estimate of the c and a





STRUCTURE (62).

FIG. IV. 10. HEXAGONAL Th2Ni17 FIG. IV. 11. SCHEMATIC REPRE-SENTATION OF STACKING OF BLOCKS IN Th2Zn17 STRUCTUR

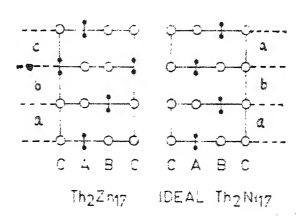


FIG. IV.12. SCHEMATIC REPRESENTATION OF Th2Zn17 - TYPE AND Th2N117 - TYPE STRUCTURES (60).

parameters was made using the hkl indices of these lines on the basis of the rhombohedral RE2Co17 structure. Since in the rhombohedral RE2Co17 structure there are three blocks, height of each block can be calculated from the determined c parameter. Assuming that the S and A phases are multiblock structures, the c/a values for structures having upto eight stacks of blocks were calculated. Indexing of the diffraction patterns of the two phases was tried with a Bunn chart at these pre-determined c/a values. The most probable lattice was determined in each case by comparing the observed d values with the d values calculated on the basis of indexing. For the A and S phases, the indexed patterns are shown in Tables IV.3 and IV.4.

The diffraction pattern of the A phase (alloy 42) could be indexed on the basis of a hexagonal cell with lattice parameters,  $a=8.495~{\rm A}^{\rm O}$ ,  $c=22.663~{\rm A}^{\rm O}$  and c/a=2.668, corresponding to a stacking of  $5\frac{1}{2}$  blocks. The indexed pattern of alloy 42 is shown in Table IV.3. The diffraction pattern of the S phase (alloy 51) could be indexed on the basis of a hexagonal cell with lattice parameters,  $a=8.551~{\rm A}^{\rm O}$ ,  $c=12.405~{\rm A}^{\rm O}$  and c/a=1.45, corresponding to a stacking of 3 blocks. The indexed pattern of alloy 51 is shown in Table IV.4. The lattice parameter of the S phase is the same as the corresponding RE $_2^{\rm Co}_{17}$  phase. However, there are extra

TABLE IV.3

X-ray Diffraction Pattern of the A-phase

Hexagonal Ccll -  $a = 8.495 \text{ }^{\circ}$ ,  $c = 22.663 \text{ }^{\circ}$  and c/a = 2.668

P THE T PROPERTY.				
Dine	Relative Intensity	hkl	<sup>d</sup> observed	<sup>d</sup> calculated
ı	VW	202	3.495	3.499
2	VW	007	3: 252	3 <b>.</b> 2 <b>3</b> 8
3	VVW	204	3.060	<b>3</b> .085
4	M	107	2.962	2.963
5	$\nabla W$	205	2.853	2.856
6	VW	116	2,805	2.823
7	M	211	2.743	2.760
8	$\nabla W$	212	2.679	2.701
9	S	009	2.520	2.518
10	S	300	2.447	2.452
11	M	215	2.373	2.370
12	VW	00,10	2.267	2,266
13	W	10,10	2.176	2,166
14	VS	305	2.152	2.157
15	vs	220	2.123	2.124
16	VVS	217	2.107	2.109
17	VS	209	2.072	2.078
18	S	00,11,306	2,058	2.060, 2.057
19	$\Psi$	11,10	1.999	1.999
.20	vs	00,12	1.890	1.890

TABLE IV.3 (contd)

Tine Mc.	Relative Intensity	h k l	dobserved	dcalculated
27	$\mathbb{W} \mathbb{V}$	219;315	1.862	1.866;1.860
22	VW	308	1.858	1.854
23	$\mathbf{W}\mathbf{V}\mathbf{V}$	229	1.630	1.624
24	VVW	410,411	1.603	1.606;1.607
25	$\nabla W$	319,325	1•584	1.585.1.582
25	M	30,12	1.496	1.496
27	M	20 ,14	1.482	1.482
28	ΛM	502	1.457	1.459
29	νw	41.7	1.436	1.438
30	$\nabla \nabla W$	00,16,330	1.416	1.4164.1.4159
31	$\nabla\nabla W$	418	1.397	1.397
32	VVW	422	1.381	1.380
33	W	33 4	1•375	1.374
34	W	507	1.341	1.340
35	VVW	00,17 <sup>x</sup>	1.333	1.333
36	VVW	336 <b>;31,</b> 13	1.325	1.326;1.325
37	VW	40,12	1.317	1.318
38	W	11,17,509	1.271	1.272;1,270
39	W	40,13	1.265	1.265
40	S	30,16	1.228	1.227
43_	S	600 <sup>x</sup>	1.226	1.226
42	W	602	1.218	1.219

TABLE IV.3 (contd.)

line Mo.	Relative Intensity	hkl	d <sub>observed</sub>	d'calculated
43	W	429;40,14	1.216	1.217;1.215
44	ន	431;11,18	1.207	1.208;1.207
45	W	20,18	1.190	1.191
46	VW	22,16	1.1786	1.1784
47	<b>W</b> V	32,14;40,15	1.1678	1.1683;1.1675
48	W	50,12	1.1616	1.1607

<sup>\*</sup>These lines were used for calculation of lattice parameters.

TABLE IV.4

X-ray Diffraction Pattern of the S-phase Hexagonal cell - a = 8.551 Å , c = 12.4045 Å and c/ $\alpha$  = 1.451

Hexag	onal cell	- a = 8.551 H	, c = 12.4045  A an	c/a = 1.451
Line No.	Relative Intensity	h k l	<sup>d</sup> observed	d calculated
1 .	VW :	201	3.538	<b>3.</b> 549
2	$\nabla W$	至	<b>3 2 2 5 5</b>	
3	$\nabla W$	202	3.164	3.180
4	ន	113	2.972	2.973
5	W	104	2.869	2.862
6	VV	210	2.810	2.799
7	$\nabla M$	<u>2</u> 11	2.717	2.731
8	$\nabla \nabla W$	×	2.618	-
9	$\nabla \mathbf{W}$	¥	2.540	-
10	VS	300	2.462	2.469
11	ន	<u>2</u> 04	2.381	2.379
12	$\mathbf{W}\mathbf{V}$	案	2.192	. <b>-</b> -
13	VS	220	22137	2.138
14	VVS	303	2.122	2.120
15	VS	214	2.084	2.079
16	M	205	2.074	2.062
17	$\nabla \mathbf{W}$	222	2.019	2.021
18	$\nabla \mathbf{W}$	106	1.988	1.993
19	$\mathbf{W}\mathbf{V}$	312	1.945	1.950
20	٧s	223	1.901	1.899
21	W	116	1.871	1.862
22	W	215	1.865	1.857
23	$\Delta M$	401_	1.829	1.831
24	VVW	402,007	1.773	1.774;1.773
25	$\nabla \nabla \mathbf{M}$	107_	1.725	1.724
26	$\nabla \mathbf{W}$	32 <b>1;</b> 216	1.674	1.683;1.664
27	$\mathbf{W}\mathbf{V}$	117	1.647	1.638
28	M	322	1.631	1.639
29	W	315;306	1.583	1.583,1.586

TABLE IV.4 (contd.)

Line No.	Relative Intensity	hkl	<sup>d</sup> observed	d calculated
30 31 32	s S V <b>W</b>	324 500 331	1.493 1.480 1.413	1.490 1.481 1.416
33	VW	503;421	1.393	1.395;1.391
34	$\nabla W$	227	1.370	1.365
35	WVV	218;109	1.358	1.357;1.356
36	M	<i>3</i> 17	1.339	1.342
37	$\nabla W$	510	1.330	1.330
38	W	326 <sup>x</sup>	1.313	1.313
39	$\nabla W$	209	1.291	1.292
40	M	505 <sub>.</sub>	1.269	1.272
41	vs	601:327	1.227	1.228;1.227
42	ន	514;10,10	1.223	1,223;1,224
43	W	431	1.216	1.212
44	W	602	1.213	1.211
45	VS	309 <sup>x</sup>	1.203	1.204
46	W	520	1.186	1.186
<b>47</b>	WV	20,10	1.177	1,177
48	WV	522	1.166	1.165
49	VW	229	1.157	1.159

These lines were used for calculation of lattice parameters

These lines could not be indexed. They are possibly due to inpurities like oxides.

diffraction lines in the 5 phase pattern which indicate that the atomic ordering in this case is different than in the RE<sub>2</sub>Co<sub>17</sub> structure. No attempt was made to determine the complete crystal structure of these phases.

The diffraction pattern of RE2Co7 phase (alloy 12) has been indexed on the basis of a hexagonal Ce2Ni7 type structure with a = 4.974 ±0, c = 24.948 A0 and c/a = 5.007. The indexed pattern of alloy 12 is shown in Table IV.5. All the diffraction lines, however, could not be indexed; there are about five diffraction lines (weak and very weak lines) which could not be indexed. These lines are not due to the other phase present in alloy 12 viz, the RECo5 phase. They may possibly be due to impurities like oxides.

Since for all the phases no single phase alloy was available, for determining lattice parameters of the A and S phases as a function of Fe content, two phase alloys with the least amount of second phase were chosen. With the help of two high angle reflections of each phase, the lattice parameters of the two phases were calculated. The calculated lattice parameters of the A and S phase alloys are tabulated in Tables IV.6(a) and (b) and are shown in Figs. IV.13(a) and (b) as a function of iron content of the alloys. As shown in the figure, the lattice parameters of the A phase alloys are practically unaffected by the addition of Fe. In the case

TABLE IV.5

X-ray Diffraction Pattern of the  $^{\Lambda}_2B_7$  - Type Phase (alloy - 12) (Hexagonal Ce $_2$ Ni $_7$  - type structure)

a = 4.974 Å

a = 24.948

c/a = 5.016

Line No.	Relative Intensity	h k 1	<sup>d</sup> observed	d calculated
1	VW.	008	3.070	3.111
2	W	106	2.986	2.996
3	W	x	2.813	en.
4	٧s	107	2.735	2.748
5	W	x	2.657	_
6	VS	108	2.525	2.527
7	$\nabla \nabla W$	00,10	2.496	2.502
8	$\nabla \nabla M$	110	2.478	2.489
9	VW	112	2.455	2.453
10	$\nabla W$	x	2.440	
11	$\nabla \nabla W$	109	2.333	2.331
12	W	114	2.313	2.321
13	$\nabla V$	00,11	2.258	2.262
14	VS	10,10	2.179	2.159
15	vvs	201	2.148	2.158
16	$\Delta M$	202	2.126	2.134
17	W	x	2.110	_
18	$\nabla W$	00,12	2.065	2.074
19	ន	204	2.043	2.046
20	M	10,11	1.997	2.005
21	W	205	1.988	1.987
22	$\nabla W$	118	1.948	1.949
23	${\tt W}{\tt V}$	x	1.929	
24	$\nabla \nabla \mathbf{M}$	10,12	1.890	1.870
25	$\nabla \nabla W$	207	1.854	1.850
26	WV	11,10	1.759	1.764

TLBLE IV.5 (contd.)

Line No.	Relative Intensity	h k 1	<sup>d</sup> observed	d calculated
27	W	10,13	1.751	1.751
28	VVW	10,14	1.641	1.645
29	VVW	212	1.626	1.624
30	$\nabla \nabla W$	213	1.589	1.607
31	$\nabla \Delta M$	11,12	1.585	1.596
32	$\mathbb{W} \nabla \mathbb{V}$	214	1.579	1.584
33	$\nabla \mathbf{W}$	00,16	1.532	1.555
34	M	20,12	1.494	1.498
35	W	10,16	1.458	1.464
<b>36</b>	$\nabla\nabla W$	11,14	1-447	1-449
37	M	300	1.439	1-444
38	M	302	1.428	1.435
39	M	20,13	1.426	1.434
40	VW	219	1.413	1.409
41	$\nabla W$	304-	1.408	1.407
42	S	20,14 <sup>×</sup>	1.373	1.373
43	S	21,10	1.370	1.368
44	$\mathbf{W}\mathbf{V}$	21,11	1.327	1.326
45	VW	20,15	1.315	1.317
46	$\nabla W$	308-	1.310	1.310
47	٧S	20,16 <sup>*</sup>	1.263	1.263

<sup>\*</sup>These lines could not be indexed. They may be due to impurities \_like oxides.

VVW : Very Very Weak

S: Strong

VW : Very Weak

VS: Very Strong

W : Weak

VVS: Very Very Strong

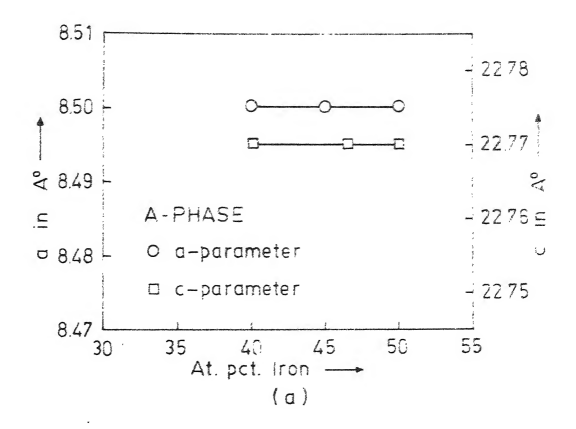
M: Medium

<sup>\*</sup>These lines were used for calculation of lattice parameters
Note: - Key to symbols used in the Tables

TABLE IV.6

Lattice Parameters of S and A phases in the RE-Co-Fe System

Alloy No.	Intended at. pct. Fe	a ( %)	° (Å)	c/a	and the second second second second
(a) <b>A</b> -p	hase		:		100
42	40.0	8 <sub>•</sub> 50	22.77	2,68	
33	46.5	8,50	22.77	2,68	
34	50.0	8.50	22.77	2.68	
, b) S-p	hase				
lı.	25 <u>.</u> 54	8,528	12.299	1.44	
25	30.0	8.547	12.340	1•44	
ī	35 <sub>•</sub> 0	8 <u>.</u> 553	12,351	1.44	
28	40.0	8,565	12,360	1.44	
51	50.0	8.572	12.365	1.44	



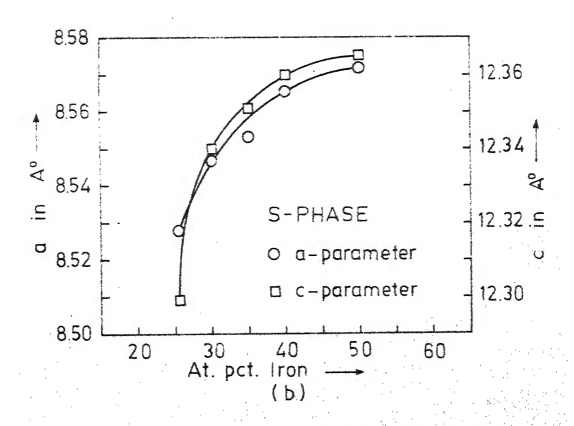


FIG. IV. 13. LATTICE PARAMETERS OF RE-Co-Fe
PHASES: (a) A-PHASE (b) S-PHASE

of the S phase, both the s and c parameters are found to increase with addition of Fe. As mentioned above, the S phase extends in a direction almost parallel to the ApB 7 Thus, the total RE content in the S phase alloys remains almost the same and the added iron is, then expected to replace the cobalt atoms in the S phase structure. Since the atomic radius of iron is not much different from that of cobalt (CN 12 atomic radii are being compared), the change in a and c parameters with replacement of Co by Fe is expected to be small. Contrary to the expected behaviour the lattice parameter data of the S phase in Fig. IV.13(b) indicate a large variation in c and a parameters with increase in Fe content; the increase being more for c than for a. observations have been reported for the RE2(Co.Fe) , 7 phases by Ray et al (59). These authors explained the abnormal caxis expansion in terms of the combined effects of increased magnetic ordering temperatures of the RE2Fe, 7 phases conferred by the addition of Co and negative exchange interactions between certain atoms, in the structures. In the present case the S phase structure is not known and whether a similar explanation for the change in lattice parameter of the S phase is applicable or not is not known. In the case of the A phase, the added Fe is expected to replace partially the RE atoms because the RE content decreases with increase in Fe content. Since

the atomic radius of Fe is substantially less than that of RE it is expected that the A phase lattice should contract as Fe is added. It is possible, however, that Fe replaces RE in pairs as in the  $^{A}_{2}^{B}_{17}$  compounds. In that case increase in c parameter is expected. Contrary to these expectations, the c and a parameters of the A phase remain unaffected on addition of Fe. In the absence of detailed structural investigations, it is difficult to explain the lattice parameter variations of the S and A phases.

### IV.2 Easy Axis of Magnetization:

In the present investigation, the easy axis of magnetization was determined by using a magnetically aligned needle of powder material and a Weissenberg camera. In taking a powder pattern with the needle formed from alloy 51 powder it was found that a good pattern, showing the streaks corresponding to preferred orientations caused by magnetic alignment of powder particles, could be obtained when the powder was etched with Nital to remove the disturbed surface layers. A similar observation has been reported for the  $Sm_2(Co,Fe)_{17}$  alloys by Sergeyev et al. (61). These authors could improve the coercivity of the  $Sm_2(Co,Fe)_{17}$  alloy powder by removing the disturbed layers from the particle surfaces by etching the powder with citric acid. The easy axis of magnetization for the etched powder of alloy 51 was found to be along the c-axis. For the other S phase alloys the easy axis has not

been determined. In the literature it has been mentioned that the addition of about 10 percent iron to the  $\mathrm{MM}_{2}\mathrm{Co}_{17}$  phase changes the easy magnetization axis from basal plane to c-axis (Fig. I.9) and this c-axis magnetic symmetry is maintained upto about 50 percent Fe (27). The earlier investigators, however, are not clear in stating the exact compositions of the alloys chosen. It appears from their report that the alloys used were along the  ${}^{\rm A}_2{}^{\rm B}_{17}$  line. It this is the case, it is possible that these authors failed to distinguish between the  ${}^{A}_{2}{}^{B}_{17}$  and the S phases. The S phase region is very close to and is almost parallel to the  $^{\Lambda}_{2}B_{17}$  line. It extends from somewhere around 20 at. pct. Fe to about 50 at. pct. Fe closely agrees with the composition range in which the c-axis magnetic symmetry has been observed in the MM-Co-Fe phase. Thus, it appears that the magnetic symmetry change is not because of Fe addition in  $^{\Lambda}_{2}$ B $_{17}$  structure but due to formation of a new phase. If this is true, then the RE2(Co,Fe)17 phase, which appears to have deviated to the higher (Co, Fe) side, should continue to show basal plane magnetic symmetry between the RE2Co17 and RE2Fe17 compositions.

## IV.3 Curie Temperature:

Induction method was adopted for determining the Curie temperatures ( $T_c$ ) of the RE-Co-Fe alloys. In the initial runs when the sample was kept inside the furnace from the beginning (i.e. during heating up of the furnace) variation

in Curie temperature was observed when the same specimen was cycled several times through the Curie temperature. indicated that the material was undergoing some transformation. To reduce the extent of transformation, the material was kept at the high temperatures for the minimum time possible. achieve this, the furnace was first heated upto 900°C (or 950°C), the temperature at which the alloys were initially annealed, and then the specimen was lowered into the furnace to heat it upto 900°C or 950°C. On reaching the temperature the specimen was drawn into the induction coil and allowed to Several traces obtained with the alleys containing different phases are shown in Figs. IV.14 to IV.17. As shown in the figures, the RE<sub>2</sub>Co., A and S phase alloys registered clear increase in induced voltages at the Chrie temperatures, But, in the case of the RE2Co., alloys large fluctuations in the induced voltage were observed. In the case of alloy 50, a three phase (RE<sub>2</sub>Co<sub>7</sub>+RECo $_{5}$ +S) alloy, two peaks in the induced voltage vs temperature traces were observed as shown in Fig. IV.16. The two temperatures correspond to the S and RECo5 phases. The change in induced voltage due to Curie temperature of the RE2Co7 phase could not be seen clearly; a small break in the trace at about 715°C (the approximate Curie temperature of the LE, Co, phase) was, however, observed. This is possibly because of the presence of only small amount.

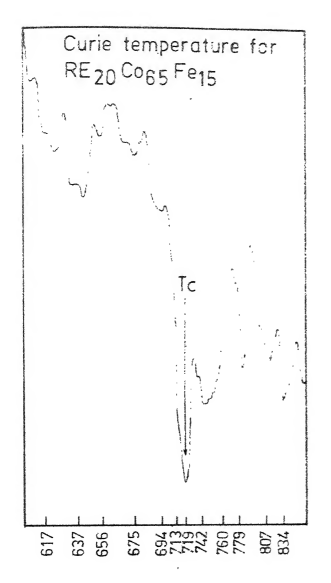
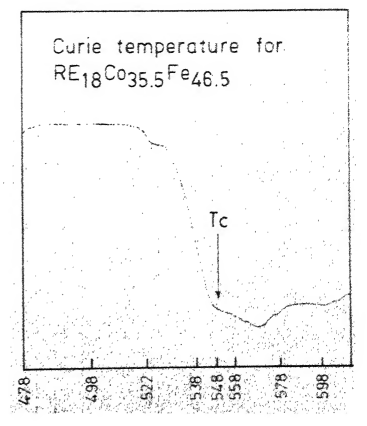
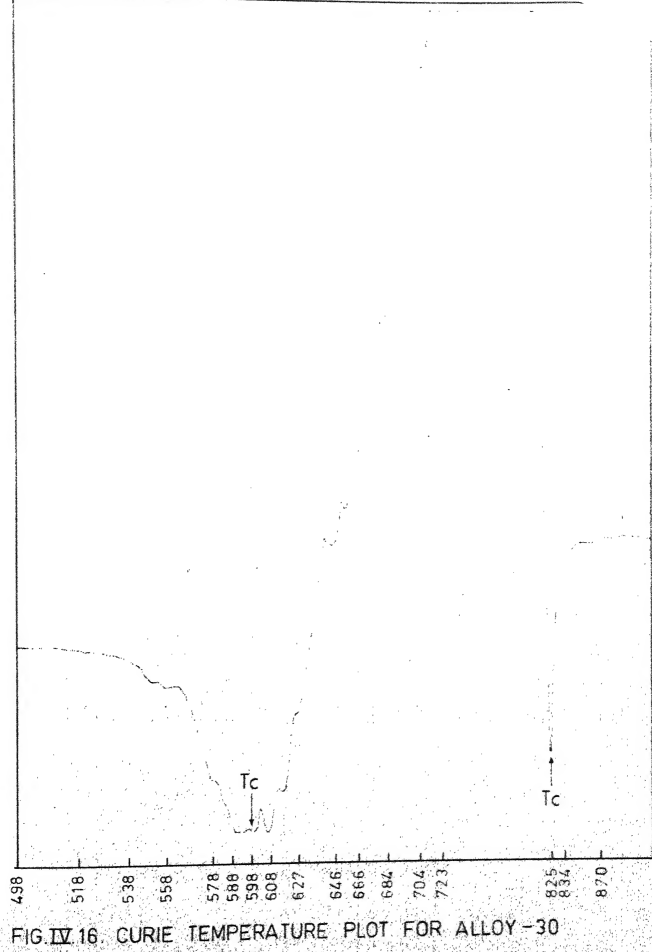


FIG IV.14. CURIE TEMPERATURE PLOT FOR RE2Ct7 PHASE.

FIG. IV.15. CURIE TEMPERATURE PLOT FOR A-PHASE.





(RE15 Co65 Fe 20 )

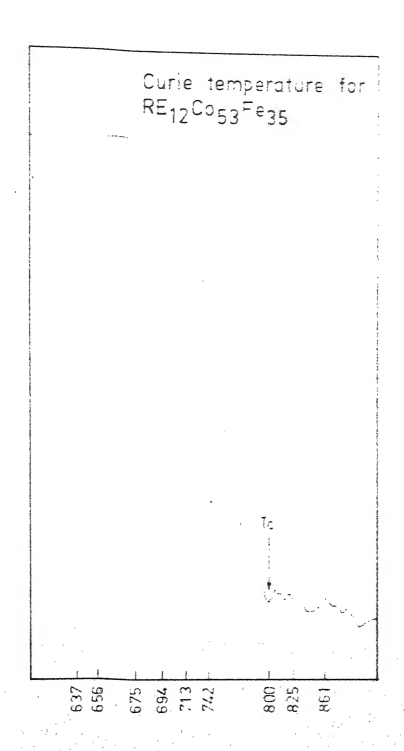


FIG. IV 17 CURIE TEMPERATURE PLOT FOR S-PHASE

of RE2Co7 phase in alloy 30. In general, a small drift was observed when the same specimen was used repeatedly for Curic temperature determination. Hence, the  $\mathbf{T}_{\mathbf{c}}$  value obtained in the first or the second run was considered to be accurate. The Curie temperatures of different alloys are tabulated in Table IV.7 and the  $\mathbb{T}_c$  values are plotted as a function of Fe content of the alloys in Fig. IV.18. As shown in the figure, the Curie temperatures of various phases appear to vary smoothly with Fe content. Among the various alloys investigated. the Curie temperatures of the A phase alloys are the lowest and those of the S phase alloys are the highest. S phase the Curie temperature appears to increase initially (upto about 25 at. pct, Fe) but with further additions of Fc, To decreases. In the case of the A phase, the Gurie temperature decreases with increase in Fe, the decrease being less marked initially. For the RE2Co7 phase the Curie temperature increases with increase in Fe. For the binary RE-Co alloys the addition of Fe is reported to decrease the Curie temperature (see Fig. I.8). But, as shown in Fig. IV.18, the Curie temperatures remain attractively high even after substantial amounts of Fe addition in the S phase.

From the data on the Curie temperatures and the easy axis of magnetization of the RE-Co-Fe alloys, it appears that the S phase offers the maximum potential for the development of permanent magnets. If the prediction that the RE<sub>2</sub>(Co,Fe)<sub>17</sub>

TABLE IV.7
Curie Temporatures of RE-Co-Fe Alloys

and the state of t	<del>,                                     </del>			
Serial No.	Alloy No.	Major Phase	Intended Compo- sition at per cent Fe	Curie Tempera- ture of the najor phase Tc, og
1				
1	12	RE2 <sup>Co</sup> 7	15	718.0
2	14	RE <sub>2</sub> Co <sub>7</sub>	20	722.5
3	52	RE <sub>2</sub> Co <sub>7</sub>	30	732.0
4	30	S-phase	20	825.0
5	48	S-phase	25	835.0
6	51.	S-phase	35	800.0
7	31	S-phase	50	700.0
8	42	Λ <b>-</b> phase	40 .	568.0
9	33	<b>Λ</b> -phase	46.5	552.0
10	. 34	A-phase	50	513.0

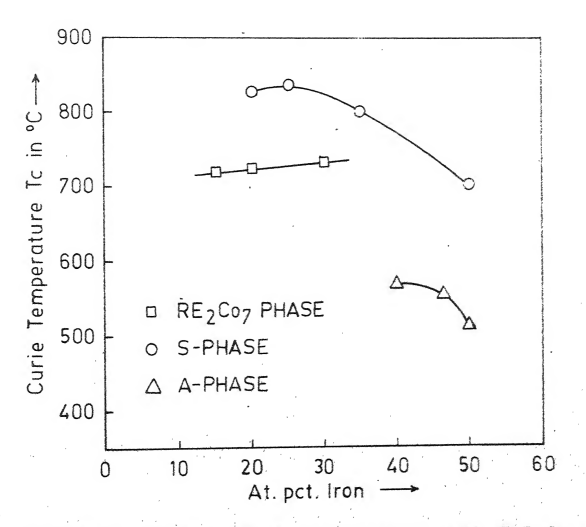


FIG. IV.18. CURIE TEMPERATURES OF RE-Co-Fe PHASES.

the S phase is true, then very high magnetization values are also expected for the S phase. Even otherwise, since the addition of Fe is reported to increase the saturation magnetization (30), it is expected that the S phase will have higher saturation magnetization values compared to the binary RECo<sub>5</sub> alloys. Thus, the S phase is expected to have better magnetic properties compared to the RECo<sub>5</sub> type and RE<sub>2</sub>Co<sub>17</sub> type phases. Also, it offers higher economic advantages compared to the RECo<sub>5</sub> and RE<sub>2</sub>Co<sub>17</sub> compounds because it takes more of iron than the RECo<sub>5</sub> type compounds and more of mischmetal than the RE<sub>2</sub>Co<sub>17</sub> type compounds.

## IV.4 Thermal Analysis:

Thermal analysis was tried with alloys which were not from the group of alloys investigated. The reason for this was that (1) due to the material used in the construction of thermal analysis apparatus, it could be used only upto 1400°C and (2) the approximate range of melting points of the investigated high Fe alloys could not be guessed. Hence, for first trial runs the alloys investigated in the present work were not suitable. Because of these reasons, two alloys with the lowest Fe content possible, 13.25 at. pct. RE, 83.8 at. pct. Co and 2.95 at. pct. Fe and 16.42 at. pct. RE, 79.92 at. pct. Co and 3.66 at. pct. Fe (57) were made use of. The usual melting points of RE-Co alloys between RECo5 and

RE2Co17 phases are in the mange of 1200°C to 1350°C. MM-Co alloys of similar compositions (but with no Fe) are also expected to show sirilar transformation temperatures. Hence, with 2 to 4 percent Fe addition (as is present in the chosen alloys) the melting points are not expected to be too far off from the binary cres, The first thermal analysis run was made with the 2.95 at. pct, Fe alloy containing RECo5 and RE<sub>2</sub>Co<sub>17</sub> phases. In this trial run the thermocouple sheath was kept outsile the alloy charge while the furnace was being heated. When the furnace temperature reached 1350°C the thermocouple along with the sheath was tried to be pushed into the melt. This was, however, not possible. Raising the furnace temperature to 1400°C also did not make it possible to insert the thermocouple into the melt. On cooling the furnace the alloy charge in the crucible was found to have nelted because the granular material charged had become a solid mass showing that melting had occured. The possible reason why the thermocouple could not be inserted in the melted mass is the presence of large amount of solid RE20017 phase having melting point higher than 1400°C. Molten part of the alloy is possibly the RECo, phase of lower melting point. Even though thermal analysis was not possible with RE-Co-Fe alloy one important observation made was that no reaction of the melt occurred with the crucible even though three hours and with the nolten alloy for more than 1/2 hr. A second trial run was made with a high RE alloy having only the RECo<sub>5</sub> phase. Since no reaction between the material charged and the alumina crucible was visible in the previous trial, in the second trial run the thermocouple along with the alumina sheath was inserted in the charge from the very beginning. The furnace was heated upto about 1250°C but at this point the furnace failed and the thermal analysis could not be carried out. On opening the thermal analysis apparatus it was found that a part of the furnace assembly had burnt out. The material charged was again found in the form of a solid mass. There was no reaction between the nelt and the crucible or the thermocouple sheath. No further work could be carried out with this apparatus.

Even though the thermal analysis work could not be carried out with the present set up the trial runs indicated that (1) the crucibles fabricated were suitable for melting as well as carrying out thermal analysis work of the RE-Co-Fe alloys, (2) the RE<sub>2</sub>Co<sub>17</sub> phase having about 3 at. pct. Fe had considerably higher melting point than the binary RE<sub>2</sub>Co<sub>17</sub> phases and (3) the melting point of RE(Co,Fe)<sub>5</sub> alloy containing about 4 at. pct. Fe had a melting point below 1250°C. The furnace assembly in the thermal analysis

apparatus worked quite woll at 1000°C and even at 1400°C during calibration and the first trial run. The temperature control and control of heating and cooling rates was quite satisfactory for carrying out thermal analysis with small amount of material.

The failure of the furnace in the second trial run is not clearly understood. Failure may be caused by exidation of Mo wire and the oxide reacting with alunina cause fusing of the refractory mass. Purified argon gas was used for protecting the furnace element. A leak in the gas purification system and supply line is a possibility. Since no abnormality was found in the gas flow, the chance of a large leak is renote. Moreover, exidation does not appear to be the real cause becaus: the charge was RE alloy and RE elements being very reactive are expected to act as a getter thereby saving Mo from oridation. The Mo wire in the sound portion of the furnace assembly also did not show any evidence of oxidation. Similarly, the RE-Co-Fe alloy charge was found to be clean and free from any oxide layer. Because of these observations, it appears that the failure of the furnace was possibly not due to oxidation. The only other possibility is the local excessive heating giving higher temperature at certain portion of the crucible and thereby melting the refractory cenent used for holding the Mo winding on the alumi former.

- 6) The easy axis of magnetization of the S phase is along the c-axis.
  - 7) The S phase, among the three phases found in this investigation, appear to offer the maximum potential for the development of permanent magnets. The S phase alloys have high Curie temperatures. The S phase also exhibits the desirable easy c-axis magnetic symmetry.
  - 8) The thermal analysis apparatus designed can be used, after making small changes, for determining the meltina points of alloys using small amounts of materials.

## SUGGESTIONS

Basing on the results obtained in the present investigation, the following suggestions can be made for full studies in this field:

- 1) The exact composition of MM can be determined in order to establish the correct composition ranges of different RE-Co-Fe phases existing in the isothermal section at 900°C.
- 2) Detailed structural investigations of the A and S phases can be carried out in order to explain the lattice parameter variations observed in these phases as a function of Fe content.
- 3) The easy axes of magnetization of the S phase alloys can be determined as a function of Fe content to ensure that the c-axis magnetic symmetry is maintained throughout the S phase region.
- 4) Even though the thermal analysis apparatus to be able to determine the melting points of the RE-Co-Fe alloys, a DTA apparatus is expected to give a more sensitive detection of melting points using still smaller amounts of charge. Since the present TA apparatus can be converted easily to a DTA apparatus, such a conversion seems to be worthwhile.
- 5) The primary magnetic properties (M $_{\rm S}$  and I $^{\rm H}c$ ) of the RE-Co-Fe alloys, particularly the S phase alloys, can be

determined in order to establish the optimum composition for developing the permanent magnets.

6) Since the S phase appears to be promising, efforman be directed at the Sabrication of magnets from the S phase alloys.

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APPENDIX I

Voltage-Gurrent-Responsition Characteristics of the Mo-wound Furnace

Room Tomperature Resistance 3.0 ohm.

Voltage volts	Current amps.	Turperature Oc
• ,		terretinestick für ihre in der eine eine eine eine eine eine eine ei
5	1.6	1.50
10	2.15	280
15	2.65	410
<b>2</b> 0	3 <u>-</u> 15	<b>5</b> 35
25	3.6	640
30	3.9	735
35	1 <sub>c*</sub> 2	815
40	4.45	890
45	4.7	960
50	4.85	1035
<b>5</b> 5	5.1	1100
60	5,3	1150
65	5 <b>.</b> 5	1200
70	5.7	1245
<b>7</b> 5	5.85	1285
80	6.0	1330
85	6.15	1370

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